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(54) Title: KUNITZ TYPE PROTEASE INHIBITORS

(57) Abstract

Analogues of the Kunitz Protease Inhibitor (KPI) domain of amyloid precursor protein bind to and inhibit activity of serine proteases, including kallikrein, plasmin and coagulation factors such as factors VIIa, DKs, Xa, XIa and XIIa. Pharmaceutical compositions containing the KPI analogues, along with methods for using such compositions, are useful for ameliorating and treating clinical conditions associated with increased serine protease activity, such as blood loss related to cardiopulmonary bypass surgery. Nucleic acid sequences encoding these analogues and systems for expression of the peptides of the invention are provided.

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KUNITZ TYPE PROTEASE INHIBITORS

Background of the Invention

The plasma, or serine, proteases of the blood contact system are known to be activated by interaction with negatively charged surfaces. For example, tissue injury during surgery exposes the vascular basement membrane, causing interaction of the blood with collagen, which is negatively charged at physiological pH. This induces a cascade of proteolytic events, leading to production of plasmin, a fibrinolytic protease, and consequent blood loss.

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Perioperative blood loss of this type can be particularly severe during cardiopulmonary bypass (CPB) surgery, in which the patient's blood flow is diverted to an artificial heart-lung machine. CPB is an essential component of a number of life-saving surgical procedures. For example, in the United States, it is estimated that 300,000 patients every year undergo coronary artery bypass grafts involving the use of CPB.

Although necessary and generally safe, CPB is associated with a significant rate of morbidity, some of which may be attributed to a "whole body inflammatory response" caused by activation of plasma protease systems and blood cells through interactions with the artificial surfaces of the heart-lung machine (Butler et al., Ann. Thorac. Surg. 55:552 (1993); Edmunds et al., J. Card. Surg. 8:404 (1993)). For example, during extracorporeal circulation, exposure of blood to negatively charged surfaces of the artificial bypass circuit, e.g., plastic surfaces in the heart-lung machine, results in direct activation of plasma factor XII.

Factor XII is a single-chain 80 kDa protein that circulates in plasma as an inactive zymogen. Contact in meganically charged mendothal all surfaces like mose the pypass direct auses surface-pound factor AII to be autoactivated to the active serine protease factor XIIa. See Colman, Agents Actions Suppl. 42:125

(1993). Surface-activated factor XIIa then processes prekallikrein (PK) to active kallikrein, which in turn cleaves more XIIa from XII in a reciprocal activation reaction that results in a rapid amplification of the contact pathway. Factor XIIa can also activate the first component of complement C1, leading to production of the anaphylatoxin C5a through the classical complement pathway.

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The CPB-induced inflammatory response includes changes in capillary permeability and interstitial fluid Cleavage of high molecular weight accumulation. kininogen (HK) by activated kallikrein generates the potent vasodilator bradykinin, which is thought to be responsible for increasing vascular permeability, resulting in edema, especially in the lung. The lung is particularly susceptible to damage associated with CPB, with some patients exhibiting what has been called "pump lung syndrome" following bypass, a condition indistinguishable from adult respiratory distress. See Johnson et al., J. Thorac. Cardiovasc. Surg. 107:1193 (1994).

Post-CPB pulmonary injury includes tissue damage thought to be mediated by neutrophil sequestration and activation in the microvasculature of the lung. (Butler et al., supra; Johnson, et al., supra). Activated factor XII can itself stimulate neutrophil aggregation. Factor XIIa-generated kallikrein, and complement protein C5a generated by Factor XIIa activation of the complement cascade, both induce neutrophil chemotaxis, aggregation and degranulation. See Edmunds et al., supra (1993). Activated neutrophils may damage tissue through release of oxygen-derived free-radicals, proteolytic enzymes such as elastase, and metabolites of arachidonic acid. Release of neutrophil products in the lung can cause changes in vascular tone, endothelial injury and loss of vascular integrity.

Intrinsic inhibition of the contact system occurs through inhibition of activated XIIa by C1-inhibitor (C1-INH). See Colman, supra. During CPB, this natural

inhibitory mechanism is overwhelmed by massive activation of plasma proteases and consumption of inhibitors. A potential therapeutic strategy for reducing post-bypass pulmonary injury mediated by neutrophil activation would, therefore, be to block the formation and activity of the neutrophil agonists kallikrein, factor XIIa, and C5a by inhibition of proteolytic activation of the contact system.

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Protease inhibitor therapy which partially attenuates the contact system is currently employed clinically in CPB. Aprotinin, also known as basic pancreatic protease inhibitor (BPPI), is a small, basic, 58 amino acid polypeptide isolated from bovine lung. It is a broad spectrum serine protease inhibitor of the Kunitz type, and was first used during bypass in an attempt to reduce the inflammatory response to CPB. See Butler et al., Aprotinin treatment results in a significant reduction in blood loss following bypass, but does not appear to significantly reduce neutrophil activation. Additionally, since aprotinin is of bovine origin, there is concern that repeated administration to patients could lead to the development of an immune response to aprotinin in the patients, precluding its further use.

The proteases inhibited by aprotinin during CPB appear to include plasma kallikrein and plasmin. (See, e.g., Scott, et al., Blood 69:1431 (1987)). Aprotinin is an inhibitor of plasmin (K, of 0.23nM), and the observed reduction in blood loss may be due to inhibition of fibrinolysis through the blocking of plasmin action. Although aprotinin inhibits plasma kallikrein, (K, of 20nM), it does not inhibit activated factor XII, and consequently only partially blocks the contact system during CPB.

Another attractive protease target for use of protease inhibitors, such as those of the present invention is factor XIIa situated at the very first

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the contact system, neutrophil activation and bradykinin release. Inhibition of XIIa would also prevent complement activation and production of C5a. More complete inhibition of the contact system during CPB could, therefore, be achieved through the use of a better XIIa inhibitor.

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Protein inhibitors of factor XIIa are known. For example, active site mutants of α_i -antitrypsin that inhibit factor XIIa have been shown to inhibit contact activation in human plasma. See Patston et al., J. Biol. Chem. 265:10786 (1990). The large size and complexity (greater than 400 amino acid residues) of these proteins present a significant challenge for recombinant protein production, since large doses will almost certainly be required during CPB. For example, although it is a potent inhibitor of both kallikrein and plasmin, nearly 1 gram of aprotinin must be infused into a patient to inhibit the massive activation of the kallikrein-kinin and fibrinolytic systems during CPB.

The use of smaller, more potent XIIa inhibitors such as the corn and pumpkin trypsin inhibitors (Wen, et al., Protein Exp. & Purif. 4:215 (1993); Pedersen, et al., J. Mol. Biol. 236:385 (1994)) could be more cost-effective than the large α_1 -antitrypsins, but the infusion of high doses of these non-mammalian inhibitors could result in immunologic reactions in patients undergoing repeat bypass operations. The ideal protein XIIa inhibitor is, therefore, preferably, small, potent, and of human sequence origin.

One candidate for an inhibitor of human origin is found in circulating isoforms of the human amyloid β -protein precursor (APPI), also known as protease nexin-2. APPI contains a Kunitz serine protease inhibitor domain known as KPI (Kunitz Protease Inhibitor). See Ponte et al., Nature, 331:525 (1988); Tanzi et al., Nature 331:528 (1988); Johnstone et al., Biochem. Biophys. Res. Commun. 163:1248 (1989); Oltersdorf et al., Nature 341:144 (1989). Human KPI shares about 45% amino acid sequence identity with aprotinin. The isolated KPI domain has

been prepared by recombinant expression in a variety of systems, and has been shown to be an active serine protease inhibitor. See, for example, Sinha, et al., J. Biol. Chem. 265:8983 (1990). The measured in vitro K, of KPI against plasma kallikrein is 45nM, compared to 20nM for aprotinin.

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Aprotinin, KPI, and other Kunitz-type serine protease inhibitors have been engineered by site-directed inhibitory activity improve mutagenesis to specificity. Thus, substitution of Lys¹⁵ of aprotinin with arginine resulted in an inhibitor with a K_i of 0.32mM toward plasma kallikrein, a 100-fold improvement over natural aprotinin. See PCT application No. 89/10374. See also Norris et al., Biol. Chem. Hoppe Seyler 371:3742 (1990). Alternatively, substitution of position 15 of aprotinin with valine or substitution of position 13 of KPI with valine resulted in elastase inhibitors with Ks in the 100 pM range, although neither native aprotinin nor native KPI significantly inhibits elastase. Wenzel et al., in: Chemistry of Peptides and Proteins, Vol. 3, (Walter de Gruyter, Berlin, New York, 1986); Sinha et al., supra. Methods for substituting residues 13, 15, 37, and 50 of KPI are shown in general terms in European Patent Application No. 0 393 431, but no specific sequences are disclosed, and no protease inhibition data are given.

Phage display methods have been recently used for preparing and screening derivatives of Kunitz-type protease inhibitors. See PCT Application No. 92/15605, which describes specific sequences for 34 derivatives of aprotinin, some of which were reportedly active as elastase and cathepsin inhibitors. The amino acid substitutions in the derivatives were distributed throughout almost all positions of the aprotinin molecule.

Phage '(qp's) perhods and also here sed to generate to carrants has bride actor list and Aassikreis see Dennis et al. ... Blos. them. 269:22129 and 269:2213 (1994). The residues that could be varied in the phage

display selection process were limited to positions 9-11, 13-17, 32, 36 and 37, and several of those residues were also held constant for each selection experiment. One of those variants was said to have a K, of 1.2nM for kallikrein, and had substitutions at positions 9 (Thr-Pro), 13 (Arg-Lys), 15 (Met-Leu), and 37 (Gly-Tyr). None of the inhibitors was tested for the ability to inhibit factor XIIa.

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It is apparent, therefore, that new protease inhibitors that can bind to and inhibit the activity of serine proteases are greatly to be desired. particular it is highly desirable to prepare peptides, based on human peptide sequences, that can inhibit proteases such as kallikrein; serine chymotrypsins A and B; trypsin; elastase; subtilisin; coagulants and procoagulants, particularly those in active form, including coagulation factors such as factors VIIa, IXa, Xa, XIa, and XIIa; plasmin; thrombin; enterokinase; acrosin; cathepsin; proteinase-3; urokinase; and tissue plasminogen activator. It is also highly desirable to prepare novel protease inhibitors that can ameliorate one or more of the undesirable clinical manifestations associated with enhanced serine protease activity, for example by reducing pulmonary damage or blood loss during CPB.

Summary of the Invention

The present invention relates to peptides that can bind to and preferably exhibit inhibition of the activity of serine proteases. Those peptides can also provide a means of ameliorating, treating or preventing clinical conditions associated with increased activity of serine proteases. Particularly, the novel peptides of the present invention preferably exhibit a more potent and specific (i.e., greater) inhibitory effect toward serine proteases of interest in comparison to known serine protease inhibitors. Examples of such proteases include: kallikrein; chymotrypsins A and B; trypsin; elastase; subtilisin; coagulants and proceagulants, particularly

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those in active form, including coagulation factors such as factors VIIa, IXa, Xa, XIa, and XIIa; plasmin; thrombin; proteinase-3; enterokinase; acrosin; cathepsin; urokinase; and tissue plasminogen activator.

In achieving the inhibition of serine protease activity, the invention provides protease inhibitors that can ameliorate one or more of the undesirable clinical manifestations associated with enhanced serine protease activity, for example, by reducing pulmonary damage or blood loss during CPB.

The present invention relates to protease inhibitors comprising the following amino acid sequences:

X1-Val-Cys-Ser-Glu-Gln-Ala-Glu-X2-Gly-X3-Cys-Arg-Ala-X4-X5-X6-X7-Trp-Tyr-Phe-Asp-Val-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-X4-Tyr-Gly-Gly-Cys-X9-X10-X11-X¹²-Asn-Asn-Phe-Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-Val-Cys-Gly-Ser-Ala-Ile,

wherein: X' is selected from Glu-Val-Val-Arg-Glu-, Asp, or Glu; X2 is selected from Thr, Val, Ile and Ser; X3 is selected from Pro and Ala; X4 is selected from Arg, Ala, Leu, Gly, or Met; X⁵ is selected from Ile, His, Leu, Lys, Ala, or Phe; X6 is selected from Ser, Ile, Pro, Phe, Tyr, Trp, Asn, Leu, His, Lys, or Glu; X' is selected from Arg, His, or Ala; X⁴ is selected from Phe, Val, Leu, or Gly; X is selected from Gly, Ala, Lys, Pro, Arg, Leu, Met, or Tyr; X10 is selected from Ala, Arg, or Gly; X11 is selected from Lys, Ala, or Asn; and X12 is selected from Ser, Ala, or Arg.

The invention relates more specifically to protease inhibitors comprising the following amino acid sequences:

X'-Val-Cys-Ser-Glu-Gln-Ala-Glu-X'-Gly-X'-Cys-Arg-Ala-X4-X5-X4-X7-Trp-Tyr-Phe-Asp-Val-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-X4-Tyr-Gly-Gly-Cys-X4-X11-X12-Asn-Asn-Phe-Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-Val-Cys-Gly-Ser-Ala-Ile,

wherein X1 is selected from Glu-Val-Val-Arg-Glu-, Asp, or

wiennes from the and and selected from AIC Geu, Gly, or Mat; X' is selected from Ile, His, Leu, Lys. Ala, or Phe; X⁶ is selected from Ser, Ile, Pro, Phe, Tyr, Trp, Asn, Leu, His, Lys, or Glu; X⁷ is selected from Arg, His, or Ala; X⁸ is selected from Phe, Val, Leu, or Gly; X⁸ is selected from Gly, Ala, Lys, Pro, Arg, Leu, Met, or Tyr; X¹⁰ is selected from Ala, Arg, or Gly; X¹¹ is selected from Lys, Ala, or Asn; X¹² is selected from Ser, Ala, or Arg; provided that when X⁴ is Arg, X⁶ is Ile; when X⁹ is Arg, X⁶ is Ala or Leu; when X⁹ is Tyr, X⁴ is Ala or X⁵ is His; and either X⁵ is not Ile; or X⁶ is not Ser; or X⁶ is not Leu, Phe, Met, Tyr, or Asn; or X¹⁰ is not Gly; or X¹¹ is not Asn; or X¹² is not Arg.

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Another aspect of this invention provides protease inhibitors wherein at least two amino acid residues selected from the group consisting of X^4 , X^5 , X^6 , and X^7 defined above differ from the residues found in the naturally occurring sequence of KPI. Another aspect of this invention provides protease inhibitors wherein X^{l} is Asp or Glu, X2 is Thr, X3 is Pro, and X12 is Ser. Yet another aspect of this invention provides protease inhibitors wherein X^i is Glu, X^2 is Thr, X^3 is Pro, X^4 is Met, X⁵ is Ile, X⁶ is Ser, X⁷ is Arg, x⁸ is Phe, X⁹ is Gly, X^{i0} is Gly, and X^{i1} is Asn. Another aspect of this invention provides protease inhibitors wherein Xi is Asp, X^2 is Thr, X^3 is Pro, X^4 is Arg, X^5 is Ile, X^6 is Ile, X^7 is Arg, x^{i} is Val, X^{0} is Arg, X^{10} is Ala, and X^{11} is Lys. Another aspect of this invention provides protease inhibitors wherein X1 is Glu-Val-Val-Arg-Glu-, X2 is Thr, X^3 is Pro, X^4 is Met, X^5 is Ile, X^4 is Ser, X^7 is Arg, x^4 is Phe, X^0 is Gly, X^{10} is Gly, X^{11} is Asn, and X^{12} is Ala. Another aspect of this invention provides protease inhibitors wherein X' is Glu-Val-Val-Arg-Glu-, X' is Thr, X3 is Pro, X4 is Met, X5 is Ile, X4 is Ser, X7 is Arg, x8 is Phe, X^{i} is Gly, X^{ij} is Gly, X^{ij} is Ala, and X^{ij} is Arg. Another aspect of this invention provides protease inhibitors wherein X' is Glu, X2 is Thr, X3 is Pro, X4 is Met, X^5 is Ile, X^6 is Ser, X^7 is Arg, x^8 is Phe, X^9 is Gly, X^{10} is Ala, X^{11} is Asn, and X^{12} is Arg. Another aspect of this invention provides protease inhibitors wherein X' is

Glu-Val-Val-Arg-Glu-, X² is Thr, X³ is Pro, X⁴ is Met, X³ is Ile, X⁴ is Ser, X¹ is Arg, x¹ is Phe, X⁴ is Gly, X¹0 is Arg, X¹1 is Asn, and X¹2 is Arg. Another aspect of this invention provides protease inhibitors wherein X¹ is Glu-Val-Arg-Glu-, X² is Thr, X³ is Pro, X⁴ is Met, X⁵ is Ile, X⁶ is Ser, X¹ is Arg, x¹ is Val, Leu, or Gly, X⁰ is Gly, X¹0 is Gly, X¹1 is Asn, and X¹2 is Arg. Another aspect of this invention provides protease inhibitors wherein X¹ is Glu-Val-Val-Arg-Glu-, X² is Thr, X³ is Pro, X⁴ is Met, X³ is Ile, X⁶ is Ser, X¹ is Ala, x¹ is Phe, X⁰ is Gly, X¹0 is Gly, X¹1 is Asn, and X¹2 is Arg. Another aspect of this invention provides protease inhibitors wherein X¹ is Glu-Val-Val-Arg-Glu-, X² is Thr, Val, or Ser, X³ is Pro, X⁴ is Ala or Leu, X³ is Ile, X⁶ is Tyr, X¹ His, X¹ is Phe, X⁵ is Gly, X¹10 is Gly, X¹10 is Gly, X¹11 is Ala, and X¹12 is Arg.

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Yet another aspect of this invention provides protease inhibitors wherein X^2 is Thr, and X^4 is Ala. Another aspect of this invention provides protease inhibitors wherein X^2 is Thr, and X^4 is Leu. Another aspect of this invention provides protease inhibitors wherein X^2 is Val, and X^4 is Ala. Another aspect of this invention provides protease inhibitors wherein X^2 is Ser, and X^4 is Ala. Another aspect of this invention provides protease inhibitors wherein X^2 is Val, and X^4 is Leu. Another aspect of this invention provides protease inhibitors wherein X^2 is Ser, and X^4 is Leu.

Yet another aspect of this invention provides protease inhibitors wherein X¹ is Glu-Val-Val-Arg-Glu-, X² is Thr, X³ is Pro, X⁴ is Leu, X⁵ is Phe, X⁶ is Lys, X⁷ is Arg, X⁶ is Phe, X⁹ is Gly, X¹⁰ is Gly, X¹¹ is Ala, and X¹² is Arg. Another aspect of this invention provides protease inhibitors wherein X¹ is Glu-Val-Val-Arg-Glu-, X² is Thr, X³ is Pro, X⁴ is Leu, X⁵ is Phe, X⁶ is Lys, X⁷ is Arg, X⁶ is Phe, X⁹ is Tyr, X¹⁰ is Gly, X¹¹ is Ala, and X¹² is Arg. Another aspect of this invention provides protease

The Let the MR Arg. She, A' is Leu, X' is Gly, X' is Aia, and X' is Arg.

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The present invention also relates to protease inhibitors comprising the following amino acid sequences:

X'-Val-Cys-Ser-Glu-Gln-Ala-Glu-Thr-Gly-Pro-Cys-X2-Ala-X3-X4-X5-X6-Trp-Tyr-Phe-Asp-Val-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-Phe-Tyr-Gly-Gly-Cys-Gly-Gly-Asn-Arg-Asn-Asn-Phe-Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-Val-Cys-Gly-Ser-Ala-Ile,

wherein: Xi is selected from Glu-Val-Val-Arg-Glu- and Asp-Val-Val-Arg-Glu-; X2 is selected from Arg and Lys; X3 is selected from Met, Arg, Ala, Leu, Ser, Val; X^4 is selected from Ile and Ala; X' is selected from Ser, Ile, Ala, Pro, Phe, Tyr, and Trp; and X is selected from Arg, Ala, His, Gln, and Thr; provided that: when X^2 is Arg, X^3 is Leu, and X4 is Ile, X5 cannot be Ser; and also provided that either X3 is not Met; or X4 is not Ile; or X5 is not Ser; or X is not Arg. Another aspect of this invention provides protease inhibitors wherein X3 is Arg or Met, and X³ is Ser or Ile. Yet another aspect of this invention provides protease inhibitors wherein X⁵ is selected from Phe, Tyr and Trp. Another aspect of this invention provides protease inhibitors wherein X3 is Ala or Leu.

A further aspect of this invention provides an isolated DNA molecule comprising a DNA sequence encoding a protease inhibitor of the invention. Another aspect of this invention provides an isolated DNA molecule comprising a DNA sequence encoding the protease inhibitor that further comprises an isolated DNA molecule operably linked to a regulatory sequence that controls expression of the coding sequence of the protease inhibitor in a host cell. Another aspect of this invention provides an isolated DNA molecule comprising a DNA sequence encoding the protease inhibitor operably linked to a regulatory sequence that controls expression of the coding sequence of the procease inhibitor in a host cell that further comprises a DNA sequence encoding a secretory signal peptide. That secretory signal peptide may preferably comprise the signal sequence of yeast alpha-mating factor. Another aspect of this invention provides a host cell transformed with any of the DNA molecules defined above. Such a host cell may preferably comprise *E. coli* or a yeast cell. When such a host cell is a yeast cell, the yeast cell may be selected from Saccharomyces cerevisiae and Pichia pastoris.

Another aspect of this invention provides a method for producing a protease inhibitor of the present invention, comprising the steps of culturing a host cell as defined above and isolating and purifying said protease inhibitor.

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A further aspect of this invention provides a pharmaceutical composition, comprising a protease inhibitor of the present invention together with a pharmaceutically acceptable sterile vehicle.

An additional aspect of this invention provides a method of treatment of a clinical condition associated with increased activity of one or more serine proteases, comprising administering to a patient suffering from said clinical condition an effective amount of a pharmaceutical composition comprising a protease inhibitor of the present invention together with a pharmaceutically acceptable sterile vehicle. That method of treatment may preferably be used to treat the clinical condition of blood loss during surgery.

Yet another aspect of this invention provides a method for inhibiting the activity of serine proteases of interest in a mammal comprising administering a therapeutically effective dose of a pharmaceutical composition comprising a protease inhibitor of the present invention together with a pharmaceutically acceptable sterile vehicle.

Another aspect of this invention provides a method for inhibiting the activity of serine proteases of interest in a mammal comprising administering a therapeutically effective dose of a pharmaceutical

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are selected from the group consisting of: kallikrein; chymotrypsins A and B; trypsin; elastase; subtilisin; coagulants and procoagulants, particularly those in active form, including coagulation factors such as factors VIIa, IXa, Xa, XIa, and XIIa; plasmin; thrombin; cathepsin; acrosin; proteinase-3; enterokinase; urokinase; and tissue plasminogen activator.

A further aspect of this invention relates to protease inhibitors comprising the following amino acid sequences:

X'-Val-Cys-Ser-Glu-Gln-Ala-Glu-Thr-Gly-Pro-Cys-Arg-Ala-X2-X3-X4-Arg-Trp-Tyr-Phe-Asp-Val-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-Phe-Tyr-Gly-Gly-Cys-X5-Gly-Asn-Arg-Asn-Asn-Phe-Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-Val-Cys-Gly-Ser-Ala-Ile,

wherein X^i is selected from Glu-Val-Val-Arg-Glu-, Asp, or Glu; X² is selected from Ala, Leu, Gly, or Met; X³ is selected from Ile, His, Leu, Lys, Ala, or Phe; X4 is selected from Ser, Ile, Pro, Phe, Tyr, Trp, Asn, Leu, His, Lys, or Glu; X5 is selected from Gly, Ala, Lys, Pro, Arg, Leu, Met, or Tyr; provided that when X¹ is Arg, X² is Ala or Leu; when X⁵ is Tyr, X² is Ala or X³ is His; and either X3 is not Ile; or X4 is not Ser; or X3 is not Leu, Phe, Met, Tyr, or Asn. Another aspect of this invention provides a protease inhibitor as defined above wherein Xi is Glu, X2 is Met, X3 is Ile, X4 is Ile, and X5 is Gly.

The invention also relates more specifically to protease inhibitors comprising the following amino acid sequences:

Glu-Val-Val-Arg-Glu-Val-Cys-Ser-Glu-Gln-Ala-Glu-Thr-Gly-Pro-Cys-Arg-Ala-Xi-X2-X3-Arg-Trp-Tyr-Phe-Asp-Val-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-Phe-Tyr-Gly-Gly-Cys-X'-Gly-Asn-Arg-Asn-Asn-Phe-Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-Val-Cys-Gly-Ser-Ala-Ile,

wherein X1 is selected from Ala, Leu, Gly, or Met; X2 is 35 selected from Ile, His, Leu, Lys, Ala, or Phe; X3 is selected from Ser, Ile, Pro, Phe, Tyr, Trp, Asn, Leu, His, Lys, or Glu; X' is selected from Gly, Arg, Leu, Met, or Tyr; provided that when X1 is Ala, X2 is Ile, His, or Leu; when X^i is Leu, X^2 is Ile or His; when X^i is Leu and X^2 is Ile, X^3 is not Ser; when X^i is Gly, X^2 is Ile; when X^4 is Arg, X^i is Ala or Leu; when X^4 is Tyr, X^i is Ala or X^2 is His; and either X^i is not Met, or X^2 is not Ile, or X^3 is not Ser, or X^4 is not Gly.

A further aspect of this invention provides a protease inhibitor as defined above wherein X¹ is Met, X³ is Ser, and X⁴ is Gly. Another aspect of this invention provides a protease inhibitor wherein X² is selected from His, Ala, Phe, Lys, and Leu. Another aspect of this invention provides a protease inhibitor wherein X² is His. Another aspect of this invention provides a protease inhibitor wherein X² is Ala. Another aspect of this invention provides a protease inhibitor wherein X² is Phe. Another aspect of this invention provides a protease inhibitor wherein X² is Lys. Another aspect of this invention provides a protease inhibitor wherein X² is Lys. Another aspect of this invention provides a protease inhibitor wherein X² is Leu. Another aspect of this invention provides a protease inhibitor wherein X³ is Met, X³ is Ile, and X⁴ is Gly.

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Yet another aspect of this invention provides a protease inhibitor wherein X³ is Ile. Another aspect of this invention provides a protease inhibitor wherein X³ is Pro. Another aspect of this invention provides a protease inhibitor wherein X³ is Phe. Another aspect of this invention provides a protease inhibitor wherein X³ is Tyr. Another aspect of this invention provides a protease inhibitor wherein X³ is Trp. Another aspect of this invention provides a protease inhibitor wherein X³ is Asn. Another aspect of this invention provides a protease inhibitor wherein X³ is Leu.

An additional aspect of this invention provides a protease inhibitor wherein X³ is Lys. Another aspect of this invention provides a protease inhibitor wherein X³ is His. Another aspect of this invention provides a rotease inhibitor wherein Y³ is 310. Another aspect of him invention provides increase inhibitor wherein X³ has. Another aspect of this invention provides a rotease inhibitor wherein Another aspect of this invention provides a spect of this invention provides a protect of this invention provides a rotease inhibitor wherein X³ is Lys.

protease inhibitor wherein X^2 is Ile. Another aspect of this invention provides a protease inhibitor wherein X^3 is Phe, and X^4 is Gly. Another aspect of this invention provides a protease inhibitor wherein X^3 is Tyr, and X^4 is Gly. Another aspect of this invention provides a protease inhibitor wherein X^3 is Trp, and X^4 is Gly.

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Yet another other aspect of this invention provides a protease inhibitor wherein X3 is Ser or Phe, and X4 is Arg or Tyr. Another aspect of this invention provides a protease inhibitor wherein X^2 is His or Leu, X^3 is Phe, and X4 is Gly. Another aspect of this invention provides a protease inhibitor wherein Xi is Leu. Another aspect of this invention provides a protease inhibitor wherein X^2 is His, X^3 is Asn or Phe, and X^4 is Gly. aspect of this invention provides a protease inhibitor wherein X^1 is Ile, X^3 is Pro, and X^4 is Gly. aspect of this invention provides a protease inhibitor wherein X^1 is Gly, X^2 is Ile, X^3 is Tyr, and X^4 is Gly. Another aspect of this invention provides a protease inhibitor wherein X^1 is Met, X^2 is His, X^3 is Ser, and X^4 is Tyr.

Additionally, another aspect of this invention relates to procease inhibitors comprising the following amino acid sequences:

X¹-Val-Cys-Ser-Glu-Gln-Ala-Glu-X²-Gly-Pro-Cys-Arg-Ala-X³-X⁴-X⁵-X⁶-Trp-Tyr-Phe-Asp-Val-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-Phe-Tyr-Gly-Gly-Cys-X⁷-Gly-Asn-Arg-Asn-Asn-Phe-Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-Val-Cys-Gly-Ser-Ala-Ile,

wherein X¹ is selected from Glu-Val-Val-Arg-Glu-, Asp, or Glu; X² is selected from Thr, Val, Ile and Ser; X³ is selected from Arg, Ala, Leu, Gly, or Met; X⁴ is selected from Ile, His, Leu, Lys, Ala, or Phe; X⁵ is selected from Ser, Ile, Pro, Phe, Tyr, Trp, Asn, Leu, His, Lys, or Glu; X⁶ is selected from Arg, His, or Ala; and X⁷ is selected from Gly, Ala, Lys, Pro, Arg, Leu, Met, or Tyr.

Another aspect of this invention provides a protease inhibitor as defined above wherein at least two amino acid residues selected from the group consisting of X^3 ,

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 X^4 , X^5 , and X^6 differ from the residues found in the naturally occurring sequence of KPI. Another aspect of this invention provides a protease inhibitor wherein X^{l} is Glu-Val-Val-Arg-Glu-, X2 is Thr, Val, or Ser, X3 is Ala or Leu, X^4 is Ile, X^5 is Tyr, X^6 is His and X^7 is Gly. Another aspect of this invention provides a protease inhibitor wherein X^2 is Thr, and X^3 is Ala. aspect of this invention provides a protease inhibitor wherein X^2 is Thr, and X^3 is Leu. Another aspect of this invention provides a protease inhibitor wherein X^2 is Val, and X3 is Ala. Another aspect of this invention provides a protease inhibitor wherein X2 is Ser, and X3 is Another aspect of this invention provides a protease inhibitor wherein X^2 is Val, and X^3 is Leu. Another aspect of this invention provides a protease inhibitor wherein X^2 is Ser, and X^3 is Leu. aspect of this invention provides a protease inhibitor wherein X¹ is Glu-Val-Val-Arg-Glu-, X² is Thr, X³ is Leu, X^4 is Phe, X^3 is Lys, X^4 is Arg and X^7 is Gly. Another aspect of this invention provides a protease inhibitor wherein X' is Glu-Val-Val-Arg-Glu-, X2 is Thr, X3 is Leu, X^4 is Phe, X^5 is Lys, X^4 is Arg and X^7 is Tyr. Another aspect of this invention provides a protease inhibitor wherein X' is Glu-Val-Val-Arg-Glu-, X2 is Thr, X3 is Leu, X^4 is Phe, X^5 is Lys, X^6 is Arg and X^7 is Leu.

Other objects, features and advantages of the present invention will become apparent from the following detailed description. It should be understood, however, that the detailed description and the specific examples, while indicating preferred embodiments of the invention, are given by way of illustration only, since various changes and modifications within the spirit and scope of the invention will become apparent to those skilled in the art from this detailed description.

Brief Description of the Drawings

Figure 2 shows the sequence of the synthetic gene for KPI $(1 \rightarrow 57)$ fused to the bacterial phoA secretory signal sequence.

Figure 3 shows the strategy for construction of plasmid pKPI-61.

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Figure 4 shows the 192 bp XbaI-HindIII synthetic gene fragment encoding KPI (1 \rightarrow 57) and four amino acids from yeast alpha-mating factor.

Figure 5 shows the synthetic 201 bp XbaI-HindIII fragment encoding KPI(-4 \Rightarrow 57) in PKPI-61.

Figure 6 shows the strategy for the construction of plasmid pTW113.

Figure 7 shows plasmid PTW113, encoding the 445 bp synthetic gene for yeast alpha-factor-KPI($-4 \rightarrow 57$) fusion.

Figure 8 shows the amino acid sequence for KPI (-4-57).

Figure 9 shows the strategy for constructing plasmid pTW6165.

Figure 10 shows plasmid, PTW6165, encoding the 445 bp synthetic gene for alpha-factor-KPI(-4+57; M15A, S17W) fusion.

Figure 11 shows the sequences of the annealed oligonucleotide pairs used to construct plasmids PTW6165, pTW6166, pTW6175, pBG028, pTW6183, pTW6184, pTW6185, pTW6173, and pTW6174.

Figure 12 shows the sequence of plasmid PTW6166 encoding the fusion of yeast alpha-factor and KPI(-4-57; ML5A, S17Y).

Figure 13 shows the sequence of plasmid PTW6175 encoding the fusion of yeast alpha-factor and KPI(-4-57; M15L, S17F).

Figure 14 shows the sequence of plasmid PBG028 encoding the fusion of yeast alpha-factor and KPI(-4+57; MI5L, S17Y).

Figure 15 shows the sequence of plasmid PTW6183 encoding the fusion of yeast alpha-factor and KPI(-4-57; I16H, S17F).

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Figure 16 shows the sequence of plasmid PTW6184 encoding the fusion of yeast alpha-factor and KPI(-4→57; I16H, S17Y).

Figure 17 shows the sequence of plasmid PTW6185 encoding the fusion of yeast alpha-factor and KPI(-4→57; I16H, S17W).

Figure 18 shows the sequence of plasmid PTW6173 encoding the fusion of yeast alpha-factor and KPI(-4→57; M15A, I16H).

Figure 19 shows the sequence of plasmid PTW6174 10 encoding the fusion of yeast alpha-factor and KPI(-4→57; M15L. I16H).

> Figure 20 shows the amino acid sequence of KPI (-4→57; M15A, S17W).

Figure 21 shows the amino acid sequence of KPI 15 (-4→57; M15A, S17Y).

> Figure 22 shows the amino acid sequence of KPI (-4→57; M15L, S17F).

Figure 23 shows the amino acid sequence of KPI 20 (-4-57; M15L, S17Y).

> Figure 24 shows the amino acid sequence of KPI (-4→57; I16H, S17F).

> Figure 25 shows the amino acid sequence of KPI (-4→57; I16H, S17Y).

Figure 26 shows the amino acid sequence of KPI 25 (-4→57; I16H, S17W).

> Figure 27 shows the amino acid sequence of KPI (-4→57; M15A, S17F).

Figure 28 shows the amino acid sequence of KPI 30 (-4→57; M15A, I16H).

Figure 29 shows the amino acid sequence of KPI (-4-57; M15L, I16H).

Figure 30 shows the construction of plasmid pSP26:Amp:F1.

35 Figure 31 shows the construction of plasmid pgIII. Figure 32 shows the construction of plasmid

Hum. Mi.

1151......... quite **ANCE** _asm1 Figure 34 shows the construction of plasmid pALS1.

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Figure 35 shows the construction of plasmid pAL53.

shows the construction of Figure 36 PSP26:Amp:F1:PhoA:KPI:gIII.

Figure 37 shows the construction of plasmid pDW1 #14.

Figure 38 shows the coding region for the fusion of phoA-KPI (1→55)-geneIII.

Figure 39 shows the construction of plasmid PDW1 14-2.

Figure 40 shows the construction of KPI Library 16-10 19.

Figure 41 shows the expression unit encoded by the members of KPI Library 16-19.

Figure 42 shows the phoA-KPI(1→55)-geneIII region encoded by the most frequently occurring randomized KPI region.

Figure 43 shows the construction of pDD185 KPI (-4→57; M15A, S17F).

Figure 44 shows the sequence of alpha-factor fused to KPI (-4-57; M15A, S17F).

20 Figure 45 shows the inhibition constants determined for purified KPI variants against the selected serine proteases kallikrein, factor Xa, and factor XIIa.

Figure 46 shows the inhibition constants (Ks) determined for KPI variants against kallikrein, plasmin, and factors Xa, XIa, and XIIa.

Figure 47 shows the post-surgical blood loss in pigs in the presence (KPI) and absence (NS) of KPI 185-1 (M15A, S17F).

Figure 48 shows the post-surgical hemoglobin loss in 30 pigs in the presence (KPI) and absence (NS) of KPI 185-1 (M15A, S17F).

Figure 49 shows the oxygen tension in the presence and absence of KPI, before CPB, immediately after CPB, and at 60 and 180 minutes after the end of CPB.

35 Figure 50 summarizes the results shown in Figures 47-49.

Figure 51 shows the sequence of plasmid PTW6166 encoding the fusion of yeast alpha-factor and KPI(-4→57; M15A, S17Y).

Figure 52 shows the sequence of plasmid PTW6175 encoding the fusion of yeast alpha-factor and KPI($-4 \rightarrow 57$; M15L, S17F).

Figure 53 shows the sequence of plasmid PBG028 encoding the fusion of yeast alpha-factor and KPI($-4 \rightarrow 57$; M15L, S17Y).

Figure 54 shows the inhibition constants (K_is) determined for KPI variants against kallikrein, plasmin, and factor XIIa.

<u>Detailed Description</u>

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The present invention provides peptides that can bind to and preferably inhibit the activity of serine proteases. These inhibitory peptides can also provide a means of ameliorating, treating or preventing clinical conditions associated with increased activity of serine proteases. The novel peptides of the present invention preferably exhibit a more potent and specific (i.e., greater) inhibitory effect toward serine proteases of interest than known serine protease inhibitors. Examples of such proteases include: kallikrein; chymotrypsins A and B; trypsin; elastase; subtilisin; coagulants and procoagulants, particularly those in active form, including coagulation factors such as factors VIIa, IXa, Xa, XIa, and XIIa; plasmin; thrombin; proteinase-3; enterokinase; acrosin; cathepsin; urokinase; and tissue plasminogen activator.

Peptides of the present invention may be used to reduce the tissue damage caused by activation of the proteases of the contact pathway of the blood during surgical procedures such as cardiopulmonary bypass (CPB). Inhibition of contact pathway proteases reduces the "whole body inflammatory response" that can accompany contact pathway activation, and that can lead to tissue damage, and possibly death. The peptides of the present invention may also be used in conjunction with surgical

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particularly severe during CPB surgery. Pharmaceutical compositions comprising the peptides of the present invention may be used in conjunction with surgery such as CPB; administration of such compositions may occur preoperatively, perioperatively or postoperatively. Examples of other clinical conditions associated with increased serine protease activity for which the peptides of the present invention may be used include: induced inflammatory response; post-CPB pulmonary injury; pancreatitis; allergy-induced protease release; deep vein thrombosis; thrombocytopenia; rheumatoid arthritis; adult respiratory distress syndrome; chronic inflammatory bowel disease; psoriasis; hyperfibrinolytic hemorrhage; organ preservation; wound healing; and myocardial infarction. Other examples of preferable uses of the peptides of the present invention are described in U.S. Patent No. 5,187,153.

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The invention is based upon the novel substitution of amino acid residues in the peptide corresponding to the naturally occurring KPI protease inhibitor domain of human amyloid β -amyloid precursor protein (APPI). These substitutions produce peptides that can bind to serine proteases and preferably exhibit an inhibition of the activity of serine proteases. The peptides also preferably exhibit a more potent and specific serine inhibition than known serine protease inhibitors. In accordance with the invention, peptides are provided that may exhibit a more potent and specific inhibition of one or more serine proteases of interest, e.g., kallikrein, plasmin and factors Xa, XIa, XIIa, and XIIa.

The present invention also includes pharmaceutical compositions comprising an effective amount of at least one of the peptides of the invention, in combination with a pharmaceutically acceptable sterile vehicle, as described in REMINGTON'S PHARMACEUTICAL SCIENCES: DRUG RECEPTORS AND RECEPTOR THEORY, (18th ed.), Mack Publishing Co., Easton, PA (1990).

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A. Selection of sequences of KPI variants

The sequence of KPI is shown in Table 1. Table 2 shows a comparison of this sequence with that of aprotinin, with which it shares about 45% sequence identity. The numbering convention for KPI shown in Table 1 and used hereinafter designates the first glutamic acid residue of KPI as residue 1. This corresponds to residue number 3 using the standard numbering convention for aprotinin.

The crystal structure for KPI complexed with trypsin has been determined. See Perona et al., J. Mol. Biol. 230:919 (1993). The three-dimensional structure reveals two binding loops within KPI that contact the protease. The first loop extends from residue Thr⁹ to Ile¹⁶, and the second loop extends from residue Phe²¹ to Gly³¹. The two protease binding loops are joined through the disulfide bridge extending from Cys¹² to Cys³⁴. KPI contains two other disulfide bridges, between Cys³ and Cys³³, and between Cys³⁴ to Cys⁴⁹.

This structure was used as a guide to inform our strategy for making the amino acid residue substitutions that will be most likely to affect the protease inhibitory properties of KPI. Our examination of the structure indicated that certain amino acid residues, including residues 9, 11, 13-18, 32, and 37-40, appear to be of particular significance in determining the protease binding properties of the KPI peptide. In a preferred embodiment of the invention two or more of those KPI peptide residues are substituted; such substitutions preferably occurring among residues 9, 11, 13-18, 32, and In particular, we found that those substituted peptides, including peptides comprising substitutions of at least two of the four residues at positions 15-18, may exhibit more potent and specific serine protease inhibition toward selected serine proteases of interest than exhibited by the natural KPI peptide domain. Such substituted peptides may further comprise one or more dditiona, audstitutions ... esiques

and 40. ... particular, son peptides may arther

comprise a substitution at positions 9 or 37, or an additional substitution at residue 13. In particular, the peptides of the present invention preferably exhibit a greater potency and specificity for inhibiting one or more serine proteases of interest (e.g., kallikrein, plasmin and factors VIIa, IXa, Xa, XIa, and XIIa) than the potency and specificity exhibited by native KPI or other known serine protease inhibitors. That greater potency and specificity may be manifested by the peptides of the present invention by exhibiting binding constants for serine proteases of interest that are less than the binding constants exhibited by native KPI, or other known serine protease inhibitors, for such proteases.

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As an initial guide to informing the choices of amino acid substitution for preparation of KPI variants, the sequences and protease inhibitory activities of aprotinin and KPI are compared. Aprotinin is twice as potent as wild-type KPI with respect to inhibition of human plasma kallikrein, and is 100-fold more potent as an inhibitor of human plasmin. There are three amino acid differences between aprotinin and wild-type KPI in the first protease binding loop extending from residues 9 to 17. A series of KPI variants may then be created, using the methods detailed below, where the residues present in aprotinin at positions 13, 15 and 17 are substituted with the residues found in KPI. The effect of such substitutions upon KPI inhibition of plasma kallikrein and plasmin is then determined.

These results show that substitution of arginine at position 13 by lysine significantly reduces the activity of the resulting protein as an inhibitor of plasma kallikrein. Similarly, substituting positions 15 and 17 of KPI with the corresponding residues found in aprotinin also decreases potency of the KPI variants against kallikrein. Substitutions of aprotinin residues at positions 13 and 15, however, increase the potency of KPI toward plasmin. The single change of methionine to arginine at position 15 (designated M15R) decreases the K, against plasmin more than 10-fold. The change of

serine to isoleucine at position 17 (S17I) decreases the potency of KPI toward plasmin.

It is observed that single-amino acid substitutions in the first protease binding loop are generally additive, that is, combinations of single amino-acid substitutions, each of which individually enhance the potency toward plasmin, result in variants with even higher potency. The substitution R13K results in a plasmin K, of 12.3, and the further exchange of M15R results in a K, that is reduced to 1.45.

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It appears, therefore, from these results that combinations of favorable single amino acid substitutions can result in enhanced potency of KPI variants. It is further apparent that substitution in KPI with the residues found in the aprotinin first protease binding loop is not always useful. Although aprotinin is a more potent kallikrein inhibitor than KPI, none of the combinations of aprotinin residues in KPI improve kallikrein inhibition.

To further investigate substitutions that might usefully enhance protease inhibition, a series of single substitutions in KPI is prepared where charged residues in the first protease binding loop are systematically replaced with alanine. This is intended to determine whether substitutions at these sites affect potency toward plasma kallikrein, factor XIIa or plasmin.

It is found that replacement of arginine at position 13 (R13A) drastically reduces KPI inhibition of kallikrein, XIIa or plasmin. The replacement I16A, however, significantly increases the K, towards both kallikrein and plasmin, suggesting that this amino acid position is critical to inhibition of these proteases. The S17A substitution has little effect. The substitution R18A has little effect upon plasmin inhibition, but significantly impacts inhibition of kallikrein and factor XIIa. These results suggest that

ignificantly water of oteno TPI sward kallikrein or plasmin.

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These results also suggest that substitutions at residues M15 and S17 could have major effects upon inhibition of kallikrein, XIIa or plasmin. To investigate this further, two sets of yeast expression plasmids are prepared, using the methods described in detail below, in which either M15 or S17 are replaced with all possible amino acids.

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Yeast are transformed with these two sets of plasmids, and 100 individual colonies are picked at random from each transformation. Small cultures are grown from each of these colonies, and their conditioned broth is harvested and tested for kallikrein inhibiting activity. The plasmids from colonies yielding cultures expressing KPI variants more potent than wild-type KPI are isolated, and the KPI domain are sequenced. It is found that only four 4 substitutions at position 15: M15A,M15L,M15S,M15V; and 4 substitutions at position 17: S17P,S17F,S17Y and S17W, result in KPI variants with improved potency toward kallikrein.

Combinations of these position 15 and 17 mutants are then prepared to test if their effects on potency of protease inhibition are additive. Four of these double mutants ([M15A,S17Y], [M15A,S17W], [M15L,S17Y] and [M15L,S17F]) are substantially more potent toward kallikrein and factor XIIa than the single amino acid substitutions on which they are based.

The results of changing arginine at positions 18 for alanine also suggest that substitutions at position 18 could affect inhibition of kallikrein and factor XIIa. The KPI double variant M13A,S17W (named TW6165 below) is used to construct a series of variants where all possible amino acid substitutions other than Cys and Arg are placed at position 18. Of these variants, three ([M13A,S17W, R18H], [M13A,S17W, R18Q], and [M13A,S17W, R18T]) are found to exhibit enhanced inhibition of kallikrein and Factor XIIa.

The results described above relate to proteins having the N-terminal sequence EVVREVCS- et seq., as found in KPI $(-4 \rightarrow 57)$. The present invention also relates, however

to proteins wherein the N-terminal sequence may be varied, preferably by substituting aspartic acid at the N-terminus in place of the glutamic acid (i.e. the N-terminal sequence is DVVREVCS-). Other N-terminal sequences that may be used will be apparent to the skilled artisan, including a sequence lacking the first four amino acids of KPI($-4\rightarrow 57$), i.e. having the sequence EVCS-.

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By way of example, and as set forth in greater detail below, the serine protease inhibitory properties of peptides of the present invention were measured for the serine proteases of interest — kallikrein, plasmin and factors Xa, XIa, and XIIa. Methodologies for measuring the inhibitory properties of the KPI variants of the present invention are known to those skilled in the art, e.g., by determining the inhibition constants of the variants toward serine proteases of interest, as described in Example 4, infra. Such studies measure the ability of the novel peptides of the present invention to bind to one or more serine proteases of interest and to preferably exhibit a greater potency and specificity for inhibiting one or more serine protease of interest than known serine protease inhibitors such as native KPI.

The ability of the peptides of the present invention to bind one or more serine proteases of interest, particularly the ability of the peptides to exhibit such greater potency and specificity toward serine proteases of interest, manifest the clinical and therapeutic applications of such peptides. The clinical and therapeutic efficacy of the peptides of the present invention can be assayed by in vitro and in vivo methodologies known to those skilled in the art, e.g., as described in Example 5, infra.

Table 1: SEQUENCE OF KPI:

40 50 PPYGGCGGNRNNPDTRBYCMAVCGSAI

Table 2: COMPARISON OF KPI AND APROTININ SEQUENCES:

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B. Methods of producing KPI variants

The peptides of the present invention can be created by synthetic techniques or recombinant techniques which employ genomic or cDNA cloning methods.

1. Production by chemical synthesis

Peptides of the present invention can be routinely synthesized using solid phase or solution phase peptide Methods of preparing relatively short peptides such as KPI by chemical synthesis are well known in the art. KPI variants could, for example be produced by solid-phase peptide synthesis techniques using commercially available equipment and reagents such as those available from Milligen (Bedford, MA) or Applied City, (Foster Biosystems - Perkin Elmer Alternatively, segments of KPI variants could be prepared by solid-phase synthesis and linked together using segment condensation methods such as those described by Dawson et al., Science 266:776 (1994). During chemical synthesis of the KPI variants, substitution of any amino acid is achieved simply by replacement of the residue that is to be substituted with a different amino acid monomer.

Production by recombinant DNA technology (a) Preparation of genes encoding KPI variants

In a preferred embodiment of the invention, KPI variants are produced by recombinant DNA technology. This requires the preparation of genes encoding each KPI Suitable genes can be variant that is to be made. oligonucleotide synthesis constructed by commercially available equipment, such as that provided by Milligen and Applied Biosystems, supra. The genes can be prepared by synthesizing the entire coding and noncoding strands, followed by annealing the two strands. alternatively the genes can be prepared by ligation of nethous ves malie: "Vntneti. .lidonuticotildes ... the ait senes encoding Al ariants are produced for

varying the nucleotides introduced at any step of the synthesis to change the amino acid sequence encoded by the gene.

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Preferably, however, KPI variants are made by site-directed mutagenesis of a gene encoding KPI. Methods of site-directed mutagenesis are well known in the art. See, for example, Ausubel et al., (eds.) CURRENT PROTOCOLS IN MOLECULAR BIOLOGY (Wiley Interscience, 1987); PROTEIN ENGINEERING (Oxender & Fox eds., A. Liss, Inc. 1987). These methods require the availability of a gene encoding KPI or a variant thereof, which can then be mutagenized by known methods to produce the desired KPI variants. In addition, linker-scanning and polymerase chain reaction ("PCR") mediated techniques can be used for purposes of mutagenesis. See PCR TECHNOLOGY (Erlich ed., Stockton Press 1989); CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, vols. 1 & 2, loc. cit.

A gene encoding KPI can be obtained by cloning the naturally occurring gene, as described for example in U.S. Patents Nos. 5,223,482 and 5,187,153, which are hereby incorporated by reference in their entireties. In particular, see columns 6-9 of U.S. Patent No. 5,187,153. See also PCT Application No. 93/09233. In a preferred embodiment of the invention a synthetic gene encoding KPI is produced by chemical synthesis, as described above. The gene may encode the 57-amino acid KPI domain shown in Table 1, or it may also encode additional N-terminal amino acids from the APPI protein sequence, such as the four amino acid sequence (Glu-Val-Val-Arg, designated residues -4 to -1) immediately preceding the KPI domain in APPI.

Production of the gene by synthesis allows the codon usage of the KPI gene to be altered to introduce convenient restriction endonuclease recognition sites, without altering the sequence of the encoded peptide. In a preferred embodiment of the invention, the synthetic KPI gene contains restriction endonuclease recognition sites that facilitate excision of DNA cassettes from the KPI gene. These cassettes can be replaced with small

synthetic oligonucleotides encoding the desired changes in the KPI peptide sequence. See Ausubel, supra.

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This method also allows the production of genes encoding KPI as a fusion peptide with one or more additional peptide or protein sequences. The DNA encoding these additional sequences is arranged in-frame sequence encoding KPI such that, translation of the gene, a fusion protein of KPI and the additional peptide or protein sequence is produced. Methods of making such fusion proteins are well known in the art. Examples of additional peptide sequences that can be encoded in the genes are secretory signal peptide sequences, such as bacterial leader sequences, for example ompA and phoA, that direct secretion of proteins to the bacterial periplasmic space. In a preferred embodiment of the invention, the additional peptide sequence is a yeast secretory signal sequence, such as α mating factor, that directs secretion of the peptide when produced in yeast.

Additional genetic regulatory sequences can also be introduced into the synthetic gene that are operably linked to the coding sequence of the gene, thereby allowing synthesis of the protein encoded by the gene when the gene is introduced into a host cell. Examples of regulatory genetic sequences that can be introduced are: promoter and enhancer sequences and transcriptional and translational control sequences. Other regulatory sequences are well known in the art. See Ausubel et al., supra, and Sambrook et al., supra.

Sequences encoding other fusion proteins and genetic elements are well known to those of skill in the art. In a preferred embodiment of the invention, the KPI sequence is prepared by ligating together synthetic oligonucleotides to produce a gene encoding an in-frame fusion protein of yeast α -mating factor with either KPI $(1 \rightarrow 57)$ or KPI $(-4 \rightarrow 57)$.

manipulating recombinant DNA techniques that are well-

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known in the art. See, for example Sambrook et al., MOLECULAR CLONING: A LABORATORY MANUAL, Second Edition, (Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY 1989), and Ausubel, supra. In a preferred embodiment of the invention the host cell used for manipulating the KPI constructs is E. coli. For example, the construct can be ligated into a cloning vector and propagated in E. coli by methods that are well known in the art. Suitable cloning vectors are described in Sambrook, supra, or are commercially available from suppliers such as Promega (Madison, WI), Stratagene (San Diego, CA) and Life Technologies (Gaithersburg, MD).

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Once a gene construct encoding KPI has been obtained, genes encoding KPI variants are obtained by manipulating the coding sequence of the construct by standard methods of site-directed mutagenesis, such as excision and replacement of small DNA cassettes, as described supra. See Ausubel, supra, and Sinha et al., supra. See also U.S. Patent 5,373,090, which is herein incorporated by reference in its entirety. See particularly, columns 4-12 of U.S. Patent 5,272,090. These genes are then used to produce the KPI variant peptides as described below.

Alternatively, KPI variants can be produced using phage display methods. See, for example, Dennis et al. supra, which is hereby incorporated by reference in its See also U.S. Patent Nos. 5,223,409 and 5,403,484, which are hereby also incorporated by In these methods, reference in their entireties. libraries of genes encoding variants of KPI are fused inframe to genes encoding surface proteins of filamentous phage, and the resulting peptides are expressed (displayed) on the surface of the phage. The phage are then screened for the ability to bind, under appropriate conditions, to serine proteases of interest immobilized Large libraries of phage can be on a solid support. used, allowing simultaneous screening of the binding properties of a large number of KPI variants. Phage that have desirable binding properties are isolated and the sequences of the genes encoding the corresponding KPI

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variants is determined. These genes are then used to produce the KPI variant peptides as described below.

(b) Expression of RPI variant peptides

Once genes encoding KPI variants have been prepared, they are inserted into an expression vector and used to produce the recombinant peptide. Suitable expression vectors and corresponding methods of recombinant proteins and peptides are well known in the art. Methods of expressing KPI peptides are described in U.S. Patent 5,187,153, columns 9-11, U.S. 5,223,482, columns 9-11, and PCT application 93/09233, pp. 49-67. See also Ausubel et al., supra, and Sambrook et al., supra. The gene can be expressed in any number of different recombinant DNA expression systems to generate large amounts of the KPI variant, which can then be purified and tested for its ability to bind to and inhibit serine proteases of interest.

Examples of expression systems known to the skilled practitioner in the art include bacteria such as E. coli, yeast such as Saccharomyces cerevisiae and Pichia pastoris, baculovirus, and mammalian expression systems such as in Cos or CHO cells. In a preferred embodiment, KPI variants are expressed in Pichia pastoris. another preferred embodiment the KPI variants are cloned into expression vectors to produce a chimeric gene encoding a fusion protein of the KPI variant with yeast The mating factor acts as a signal α-mating factor. sequence to direct secretion of the fusion protein from the yeast cell, and is then cleaved from the fusion protein by a membrane-bound protease during the secretion process. The expression vector is transformed into S. cerevisiae, the transformed yeast cells are cultured by standard methods, and the KPI variant is purified from the yeast growth medium.

Recombinant bacterial cells expressing the peptides

... Any consummer of the recombinant antigen induced by

adding IPTG to the media or switching incubation to a higher temperature. After culturing the bacteria for a further period of between 2 and 24 hours, the cells are collected by centrifugation and washed to remove residual media. The bacterial cells are then lysed, for example, by disruption in a cell homogenizer and centrifuged to separate dense inclusion bodies and cell membranes from the soluble cell components. This centrifugation can be performed under conditions whereby dense inclusion bodies are selectively enriched by incorporation of sugars such as sucrose into the buffer and centrifugation at a selective speed. If the recombinant peptide is expressed in inclusion bodies, as is the case in many instances, these can be washed in any of several solutions to assist in the removal of any contaminating host proteins, then solubilized in solutions containing high concentrations of urea (e.g., 8M) or chaotropic agents such as guanidine hydrochloride in the presence of reducing agents such as β -mercaptoethanol or DTT (dithiothreitol).

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At this stage it may be advantageous to incubate the peptides of the present invention for several hours under conditions suitable for the peptides to undergo a refolding process into a conformation which more closely resembles that of native KPI. Such conditions generally include low protein concentrations less than 500 μ g/ml, low levels of reducing agent, concentrations of urea less than 2M and often the presence of reagents such as a mixture of reduced and oxidized glutathione which facilitate the interchange of disulphide bonds within the protein molecule. The refolding process can be monitored, for example, by SDS-PAGE or with antibodies which are specific for the native molecule (which can be obtained from animals vaccinated with the native molecule isolated from parasites). Following refolding, the peptide can then be purified further and separated from the refolding mixture by chromatography on any of several supports including ion exchange resins, gel permeation resins or on a variety of affinity columns.

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Purification of KPI variants can be achieved by standard methods of protein purification, e.g., using methods chromatographic including various liquid chromatography and adsorption performance The purity and the quality of the chromatography. peptides can be confirmed by amino acid analyses, molecular weight determination, sequence determination See, for example, PROTEIN and mass spectrometry. PURIFICATION METHODS — A PRACTICAL APPROACH, Harris et al., eds. (IRL Press, Oxford, 1989). In a preferred embodiment, the yeast cells are removed from the growth medium by filtration or centrifugation, and the KPI variant is purified by affinity chromatography on a column of trypsin-agarose, followed by reversed-phase HPLC.

C. Measurement of protease inhibitory properties of EPI variants

Once KPI variants have been purified, they are tested for their ability to bind to and inhibit serine proteases of interest in vitro. The peptides of the present invention preferably exhibit a more potent and specific inhibition of serine proteases of interest than known serine protease inhibitors, such as the natural KPI peptide domain. Such binding and inhibition can be assayed for by determining the inhibition constants for the peptides of the present invention toward serine proteases of interest and comparing those constants with for known constants determined serine inhibitors, e.g., the native KPI domain, toward those protesses. Methods for determining inhibition constants of protease inhibitors are well known in the art. See Fersht, ENZYME STRUCTURE AND MECHANISM, 2nd ed., W.H. Freeman and Co., New York, (1985).

In a preferred embodiment the inhibition experiments are carried out using a chromogenic synthetic protease substrate as described for example in Bender et al.

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constants (K values) of the peptides of the present invention toward serine proteases of interest. See Bieth in BAYER-SYMPOSIUM V "PROTEINASE INHIBITORS", Fritz et al., eds., pp. 463-69, Springer-Verlag, Berlin, Heidelberg, New York, (1974). KPI variants that exhibit potent and specific inhibition of one or more serine proteases of interest may subsequently be tested in vivo. In vitro testing, however, is not a prerequisite for in vivo studies of the peptides of the present invention.

D. Testing of KPI variants in vivo

The peptides of the present invention may be tested, alone or in combination, for their therapeutic efficacy by various in vivo methodologies known to those skilled in the art, e.g., the ability of KPI variants to reduce postoperative bleeding can be tested in standard animal models. For example, cardiopulmonary bypass surgery can be carried out on animals such as pigs in the presence of KPI variants, or in control animals where the KPI variant is not used. The use of pigs as a model for studying the clinical effects associated with CPB has previously been described. See Redmond et al., Ann. Thorac. Surg. 56:474 (1993).

The KPI variant is supplied to the animals in a pharmaceutical sterile vehicle by methods known in the art, for example by continuous intravenous infusion. Chest tubes can be used to collect shed blood for a defined period of time. The shed blood, together with the residual intrathoracic blood found after sacrifice of the animal can be used to calculate hemoglobin (Hgb) loss. The postoperative blood and Hgb loss is then compared between the test and control animals to determine the effect of the KPI variants.

E. Therapeutic use of KPI variants

EPI variants of the present invention found to exhibit therapeutic efficacy (e.g., reduction of blood loss following surgery in animal models) may preferably be used and administered, alone or in combination or as

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a fusion protein, in a manner analogous to that currently used for aprotinin or other known serine protease inhibitors. See Butler et al., supra. Peptides of the present invention generally may be administered in the manner that natural peptides are administered. therapeutically effective dose of the peptides of the present invention preferably affects the activity of the serine proteases of interest such that the clinical condition may be treated, ameliorated or prevented. Therapeutically effective dosages of the peptides of the present invention can be determined by those skilled in the art, e.g., through in vivo or in vitro models. Generally, the peptides of the present invention may be administered in total amounts of approximately 0.01 to approximately 500, specifically 0.1 to 100 mg/kg body weight, if desired in the form of one or more administrations, to achieve therapeutic effect. however, be necessary to deviate from such administration amounts, in particular depending on the nature and body weight of the individual to be treated, the nature of the medical condition to be treated, the type of preparation and the administration of the peptide, and the time interval over which such administration occurs. Thus, it may in some cases be sufficient to use less than the above amount of the peptides of the present invention, while in other cases the above amount is preferably exceeded. The optimal dose required in each case and the type of administration of the peptides of the present invention can be determined by one skilled in the art in the such view of circumstances surrounding Such peptides can be administered by administration. intravenous injections, in situ injections, applications, inhalation, oral administration using coated polymers, dermal patches or other appropriate means. Compositions comprising peptides of the present invention are advantageously administered in the form of LCL Deptides may de preferat. nectable compositions patients la continuous intravenous infusion, but can also be administered by single or

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A typical composition for such multiple injections. purpose comprises a pharmaceutically acceptable carrier. Pharmaceutically acceptable carriers include aqueous non-toxic excipients, including solutions, preservatives, buffers and the like, as described in REMINGTON'S PHARMACEUTICAL SCIENCES, pp. 1405-12 and 1461-87 (1975) and THE NATIONAL FORMULARY XIV., 14th Ed. Washington: American Pharmaceutical Association (1975). Aqueous carriers include water, alcoholic/aqueous solutions, saline solutions, parenteral vehicles such as sodium chloride, Ringer's dextrose, etc. Intravenous vehicles include fluid and nutrient replenishers. Preservatives include antimicrobials, anti-oxidants, chelating agents and inert gases. The pH and exact concentration of the various components of the composition are adjusted according to routine skills in the art. See GOODMAN AND GILMAN'S THE PHARMACOLOGICAL BASIS FOR THERAPEUTICS (7th ed.). The peptides of the present invention may be present in such pharmaceutical preparations in a concentration of approximately 0.1 to 99.5% by weight, specifically 0.5 to 95% by weight, Such pharmaceutical relative to the total mixture. preparations may also comprise other pharmaceutically active substances in addition to the peptides of the Other methods of delivering the present invention. peptides to patients will be readily apparent to the skilled artisan.

Examples of mammalian serine proteases that may exhibit inhibition by the peptides of the present invention include: kallikrein; chymotrypsins A and B; coagulants subtilisin; elastase; trypsin; procoagulants, particularly those in active form, including coagulation factors such as thrombin and and XIIa; Xa, XIa, plasmin; IXA, factors VIIa. enterokinase; acrosin: cathepsin; proteinase-3; urokinase; and tissue plasminogen activator. Examples of conditions associated with increased serine protease activity include: CPB-induced inflammatory response; post-CPB pulmonary injury; pancreatitis; allergy-induced

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protease release; deep vein thrombosis; thrombocytopenia; arthritis; adult respiratory rheumatoid syndrome; chronic inflammatory bowel disease; psoriasis; hyperfibrinolytic hemorrhage; organ preservation; wound healing; and myocardial infarction. Other examples of the use of the peptides of the present invention are described in U.S. Patent No. 5,187,153.

The inhibitors of the present invention may also be used for inhibition of serine protease activity in vitro, for example during the preparation of cellular extracts to prevent degradation of cellular proteins. For this purpose the inhibitors of the present invention may preferably be used in a manner analogous to the way that aprotinin, or other known serine protease inhibitors, are used. The use of aprotinin as a protease inhibitor for preparation of cellular extracts is well known in the art, and aprotinin is sold commercially for this purpose.

The present invention, thus generally described, will be understood more readily by reference to the following examples, which are provided by way of illustration and are not intended to be limiting of the present invention.

EXAMPLES

Example 1. Expression of wild-type KPI (-4-57)

A. Construction of PTW10:KPI

Plasmid PTW10:KPI is a bacterial expression vector encoding the 57 amino acid form of KPI fused to the bacterial phoA signal sequence. The strategy for the construction of PTW10: KPI is shown in Figure 1.

Plasmid pcDNAII (Invitrogen, San Diego, CA) was digested with PvuII and the larger of the two resulting PvuII fragments (3013 bp) was isolated. Bacterial expression plasmid pSP26 was digested with MluI and RsrII, and the 409 bp MluI-RsrII fragment containing the pTrp promoter element and transcription termination signals was isolated by electrophoresis in a 3% NuSieve طه سندست

Harose ic ML CIL untailing of leparin clamin, EGT (xe stowt) factor EGF) insert between the Ndel and Hindill sites, is

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described as pNA28 in Thompson et al., J. Biol. Chem. 269:2541 (1994). Plasmid pSP26 was deposited in host E. coli W3110, pSP26 with the American Type Culture Collection (ATCC), 12301 Parklawn Drive, Rockville, Maryland, 20852, USA under the conditions specified by the Budapest Treaty on the International Recognition of the Deposit of Microorganisms (Budapest Treaty). Host E. coli W3110, pSP26 was deposited on 3 May 1995 and given Availability of the deposited Accession No. 69800. plasmid is not to be construed as a license to practice the invention in contravention of the rights granted under the authority of any government in accordance with its patent laws.

The ends of the MluI-RsrII fragment were blunted using DNA polymerase Klenow fragment by standard The blunted fragment of pSP26 was then techniques. ligated into the large PvuII fragment of plasmid pCDNAII, and the ligation mixture was used to transform B. coli Ampicillin-resistant colonies were strain MC1061. selected and used to isolate plasmid pTW10 by standard techniques.

A synthetic gene was constructed encoding the bacterial phoA secretory signal sequence fused to the amino terminus of KPI(1-57). The synthetic gene contains cohesive ends for NdeI and HindIII, and also incorporates restriction endonuclease recognition sites for AgeI, REFII, AatII and BanHI, as shown in Figure 2. synthetic phoA-KPI gene was constructed from 6 oligonucleotides of the following sequences (shown 5'-3'):

6167: TATGAAACAAAGCACTATTGCACTGGCACTCTTACCGTTACTGTTTACCC CTGTGACAAAAGCCGAGGTGTGCTCTGAA

6169: CTCGGCTTTTGTCACAGGGGTAAACAGTAACGGTAAGAGTGCCAGTGCAA 35 TAGTGCTTTGTTTCATA

6165: CAAGCTGAGACCGGTCCGTGCCGTGCAATGATCTCCCGCTGGTACTTTGA CGTCACTGAAGGTAAGTGCGCTCCATTCTTT

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6166:
GCACTTACCTTCAGTGACGTCAAAGTACCAGCGGGAGATCATTGCACGGC
ACGGACCGGTCTCAGCTTGTTCAGAGCACAC

6168:
TACGGCGGTTGCGGCGCAACCGTAACAACTTTGACACTGAAGAGTACTG
CATGGCAGTGTGCGGATCCGCTATTTAAGCT

6164:
AGCTTAARTAGCGGATCCGCACACTGCCATGCAGTACTCTTCAGTGTCAA
AGTTGTTACGGTTGCCGCCGCAACCGCCGTAAAAGAATGGAGC

The oligonucleotides were phosphorylated and annealed 6167 + 6169, 6165 + 6166, 6168 + 6164. in pairs: 20 μl T4 DNA Ligase Buffer (New England Biolabs, Beverley, MA), 1 µg of each oligonucleotide pair was incubated with 10 U T4 Polynucleotide Kinase (New England Biolabs) for 1 h at 37°C, then heated to 95°C for 1 minute, and slow-cooled to room temperature to allow All three annealed oligo pairs were then annealing. mixed for ligation to one another in a total volume of 100 ul T4 DNA Ligase Buffer, and incubated with 400 U T4 DNA Ligase (New England Biolabs) overnight at 15°C. The ligation mixture was extracted with an equal volume of phenol: CHCl, (1:1), ethanol-precipitated, resuspended in 50 μ l Restriction Endonuclease Buffer #4 (New England Biolabs) and digested with NdeI and HindIII. annealed, ligated and digested oligos were then subjected to electrophoresis in a 3% NuSieve Agarose gel, and the 240 bp NdeI-HindIII fragment was excised. purified synthetic gene was ligated into plasmid pTW10 which had previously been digested with NdeI and HindIII, and the ligation mixture was used to transform E. coli strain MC1061. Ampicillin-resistant colonies were selected and used to prepare plasmid pTW10:KPI. plasmid contains the phoA-KPI(1-57) fusion protein inserted between the pTrp promoter element and the transcription termination signals.

The strategy for constructing OKF shows Figure 5 Plasmid pTW10:KPI was digested with Agel and

HindIII; the resulting 152 bp AgeI-HindIII fragment containing a portion of the KPI synthetic gene was isolated by preparative gel electrophoresis. An oligonucleotide pair (129 + 130) encoding the 9 aminoterminal residues of KPI(1 \rightarrow 57) and 4 amino acids of yeast α -mating factor was phosphorylated and annealed as described above.

129: CTAGATAAAAGAGAGGTGTGCTCTGAACAAGCTGAGA

130: CCGGTCTCAGCTTGTTCAGAGCACACCTCTCTTTTAT

10 The annealed oligonucleotides were then ligated to the AgeI-HindIII fragment of the KPI (1-57) synthetic gene. The resulting 192 bp XbaI-HindIII synthetic gene (shown in Figure 4) was purified by preparative gel electrophoresis, and ligated into plasmid pUC19 which had 15 previously been digested with XbaI and HindIII. ligation products were used to transform E. coli strain MC1061. Ampicillin-resistant colonies were picked and used to prepare plasmid PKPI-57 by standard methods. To create a synthetic gene encoding KPI(-4-57), PKPI-57 was 20 digested with XbsI and AgeI and the smaller fragment replaced with annealed oligos 234 + 235, which encode 4 amino acid residues of yeast α -mating factor fused a 4 amino acid residue amino-terminal extension of KPI(1→57).

234: CTAGATAAAAGAGAGGTTGTTAGAGAGGTGTGCTCTGAACAAGCTGAGA 235: CCGGTCTCAGCTTGTTCAGAGCACCTCTCTTTAT

The 4 extra amino acids are encoded in the amyloid β -protein precursor/protease nexin-2 (APPI) which contains the KPI domain. The synthetic 201 bp XbaI-HindIII fragment encoding KPI(-4+57) in pKPI-61 is shown in Figure 5.

C. Assembly of pTW113

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The strategy for the construction of PTW113 is shown in Figure 6. Plasmid pSP35 was constructed from yeast expression plasmid pYES2 (Invitrogen, San Diego, CA) as

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follows. A 267 bp PvuII-XbaI fragment was generated by PCR from yeast α -mating factor DNA using oligos 6274 and 6273:

6274: GGGGGCAGCTGTATAAACGATTAAAA

5 6273: GGGGGTCTAGAGATACCCCTTCTTCTTTAG

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This PCR fragment, encoding an 82 amino acid portion of yeast α -mating factor, including the secretory signal peptide and pro-region, was inserted into pYES2 that had been previously digested with PvuII and XbaI. The resulting plasmid is denoted pSP34.

Two oligonucleotide pairs, 6294 + 6292 were then ligated to 6290 + 6291, and the resulting 135 bp fragment was purified by gel electrophoresis.

6294: CTAGATAAAAGAGAGGCTGAGGCTCACGCTGAAGGTACTTTCACTTC

15 6290: TGACGTCTTCTTACTTGGAAGGTCAAGCTGCTAAGGAATTCAT
CGCTTGGTTGGTCAAAGGTAGAGGTTAAGCTTA

6291: CTAGTAACCTTACCTTTGACCAACCAAGCGATGAATTC

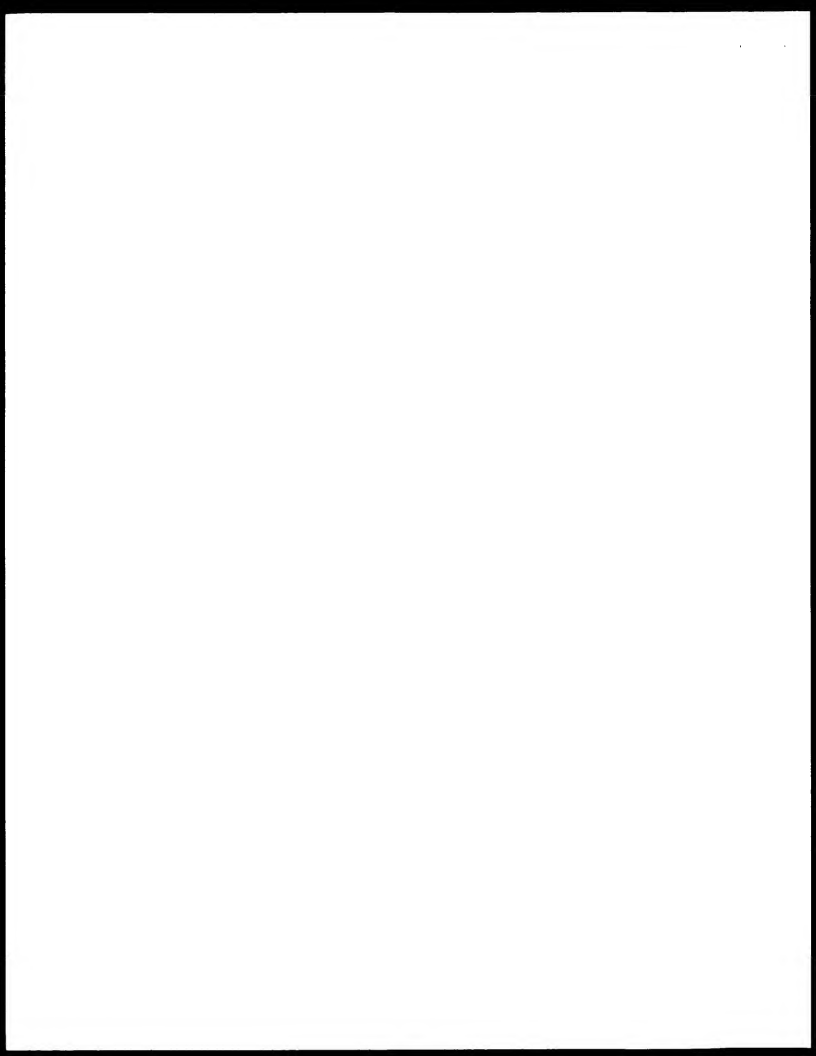
6292: GCTTGACCTTCCAAGTAAGAAGAGACGTCAGAAGTGAAAGTACCT
TCAGCGTGAGCCTCAGCCTCTTTTTAT

The resulting synthetic fragment was ligated into the XbaI site of pSP34, resulting in plasmid pSP35. pSP35 was digested with XbaI and HindIII to remove the insert, and ligated with the 201 bp XbaI-HindIII fragment of pKPI-61, encoding KPI(-4 \rightarrow 57). The resulting plasmid pTW113, encodes the 445 bp synthetic gene for the α -factor-KPI(-4 \rightarrow 57) fusion. See Figure 7.

D. Transformation of yeast with pTW113

Saccharomyces cerevisiae strain ABL115 was transformed with plasmid pTW113 by electroporation by the method of Becker et al. Methods Enzymol 194:182 (1991)

inocurate 200 Mg /PD medium. The inocurated curvaire was grown with vigorous shaking at 30°C to an OD. Of 1.3-1.5,



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at which time the cells were harvested by centrifugation at 5000 rpm for 5 minutes. The cell pellet was resuspended in 200 ml ice-cold water, respun, resuspended in 100 ml ice-cold water, then pelleted again. washed cell pellet was resuspended in 10 ml ice-cold 1M sorbitol, recentrifuged, then resuspended in a final volume of 0.2 ml ice-cold 1M sorbitol. A 40 μ l aliquot of cells was placed into the chamber of a cold 0.2 cm electroporation cuvette (Invitrogen), along with 100 ng plasmid DNA for pTW113. The cuvette was placed into an Invitrogen Electroporator II and pulsed at 1500 V, 25 μ F, 100 Ω . Electroporated cells were diluted with 0.5 ml 1M sorbitol, and 0.25 ml was spread on an SD agar plate containing 1M sorbitol. After 3 days' growth at 30°C, individual colonies were streaked on SD + CAA agar plates.

B. Induction of pTWll3/ABL115, purification of KPI(-4+57)

Yeast cultures were grown in a rich broth and the galactose promoter of the KPI expression vector induced with the addition of galactose as described by Sherman, Methods Enzymol. 194:3 (1991). A single well-isolated colony of pTW113/ABL115 was used to inoculate a 10 ml overnight culture in Yeast Batch Medium. The next day, 1L Yeast Batch Medium which had been made 0.2% glucose was inoculated to an ODem of 0.1 with the overnight Following 24 hours at 30°C with vigorous shaking, the 1L culture was induced by the addition of 20 ml Yeast Galactose Feed Medium. Following induction, the culture was fed every 12 hours with the addition of 20 ml Yeast Galactose Feed Medium. At 48 hours after induction, the yeast broth VAS harvested centrifugation, then adjusted to pH 7.0 with 2M Tris, pH The broth was subjected to trypsin-Sepharose affinity chromatography, and bound KPI(-4-57) was eluted The normal mode was not See Schilling at at Sene

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accompaished by HPLC chromatography on a semi-prep Vydac

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C4 column in a gradient of 20% to 35% acetonitrile. The sample was dried and resuspended in PBS at 1-2 mg/ml. The amino acid sequence of KPI(-4 \rightarrow 57) is shown in Figure 8.

5 Example 2. Recombinant Expression of site-directed KPI(-4+57) variants

Expression vectors for the production of specific variants of KPI(-4→57) were all constructed using the pTW113 backbone as a starting point. For each KPI variant, an expression construct was created by replacing the 40 bp RsrII-AatII fragment of the synthetic KPI gene of contained in pTW113 with a pair annealed oligonucleotides which encode specific codons mutated from the wild-type KPI(-4-57) sequence. In the following Examples the convention used for designating the amino substituents in the KPI variants indicates first the single letter code for the amino acid found in wild-type KPI, followed by the position of the residue using the numbering convention described supra, followed by the code for the replacement amino acid. Thus, for example, M15R indicates that the methionine residue at position 15 is replaced by an arginine.

A. Construction of pTW6165

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The strategy for constructing pTW6165 is shown in Figure 9. Plasmid pTW113 was digested with RsrII and AatII, and the larger of the two resulting fragments was isolated. An oligonucleotide pair (812 + 813) was phosphorylated, annealed and gel-purified as described above.

30 812: GTCCGTGCCGTGCAGCTATCTGGCGCTGGTACTTTGACGT

813: CARAGTACCAGCGCCAGATAGCTGCACGGCACG

The annealed oligonucleotides were ligated into the RsrII and AatII-digested pTW113, and the ligation product was used to transform *B. coli* strain MC1061. Transformed colonies were selected by ampicillin resistance. The

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resulting plasmid, pTW6165, encodes the 445 bp synthetic gene for the α -factor-KPI(-4 \rightarrow 57; M15A, S17W) fusion. See Figure 10.

B. Construction of pTW6166, pTW6175, pBG028, pTW6183, pTW6184, pTW6185, pTW6173, pTW6174.

Construction of the following KPI($-4 \rightarrow 57$) variants was accomplished exactly as outlined for pTW6165. The oligonucleotides utilized for each construct are denoted below, and the sequences of annealed oligonucleotide pairs are shown in Figure 11. Figures 12-19 show the synthetic genes for the α -factor fusions with each KPI($-4 \rightarrow 57$) variant.

pTW6166: KPI(-4+57; M15A, S17Y) - See Figure 12

814: GTCCGTGCCGTGCAGCTATCTACCGCTGGTACTTTGACGT

15 815: CAAAGTACCAGCGGTAGATAGCTGCACGGCACG

pTW6175: KPI(-4-57; M15L, S17F) — See Figure 13

867: GTCCGTGCCGTGCATTGATCTTCCGCTGGTACTTTGACGT

868: CAAAGTACCAGCGGAAGATCAATGCACGGCACG

pBG028: KPI(-4+57; M15L, S17Y) - See Figure 14

20 1493: GTCCGTGCCGTGCTTTGATCTACCGCTGGTACTTTGACGT

1494: CARAGTACCAGCGGTAGATCARAGCACGGCACG

pTW6183: KPI(-4-57; I16H, S17F) — See Figure 15

925: GTCCGTGCCGTGCAATGCACTTCCGCTGGTACTTTGACGT

926: CARAGTACCAGCGGAAGTGCATTGCACGGCACG

127: JTCCGTGCCGTGCAATGCACTACCGCTGGTACTTTGACGT

928: CAAAGTACCAGCGGTAGTGCATTGCACGGCACG

pTW6185: KPI(-4-57; I16H, S17W) - See Figure 17

929: GTCCGTGCCGTGCAATGCACTGGCGCTGGTACTTTGACGT

930: CAAAGTACCAGCGCCAGTGCATTGCACGGCACG

5 pTW6173: KPI(-4→57; M15A, I16H) — See Figure 18

863: GTCCGTGCCGTGCAGCTCACTCCCGCTGGTACTTTGACGT

864: CAAAGTACCAGCGGGAGTGAGCTGCACGGCACG

pTW6174: KPI(-4→57; M15L, I16H) — See Figure 19

865: GTCCGTGCCGTGCATTGCACTCCCGCTGGTACTTTGACGT

10 866: CAAAGTACCAGCGGGAGTGCAATGCACGGCACG

- C. Transformation of yeast with expression vectors Yeast strain ABL115 was transformed by electroporation exactly according to the protocol described for transformation by pTW113.
- D. Induction of transformed yeast strains, purification of KPI(-4-57) variants.

Cultures of yeast strains were grown and induced, and recombinant secreted KPI($-4 \rightarrow 57$) variants were purified according to the procedure described for KPI($-4 \rightarrow 57$). The amino acid sequences of KPI($-4 \rightarrow 57$) variants are shown in Figures 20-29.

- Example 3. Identification of KPI (-4-57; M15A, S17F) DD185 by phage display.
 - A. Construction of vector pSP26:Amp:F1
- The construction of pSP26:Amp:F1 is outlined in Figure 30. Vector pSP26:Amp:F1 contributes the basic plasmid backbone for the construction of the phage display vector for the phoA:KPI fusion, PDW1 #14. pSP26:Amp:F1 contains a low-copy number origin of

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replication, the ampicillin-resistance gene (Amp) and the F1 origin for production of single-stranded phagemid DNA.

The ampicillin-resistance gene (Amp) was generated through polymerase chain reaction (PCR) amplification from the plasmid genome of PUC19 using oligonucleotides 176 and 177.

176: GCCATCGATGGTTTCTTAAGCGTCAGGTGGCACTTTTC

177: GCGCCAATTCTTGGTCTACGGGGTCTGACGCTCAGTGGAACGAA

The PCR amplification of Amp was done according to standard techniques, using Taq polymerase (Perkin-Elmer Cetus, Norwalk, CT). Amplification from plasmid pUC19 with these oligonucleotides yielded a fragment of 1159 bp, containing PflMI and ClaI restriction sites. The PCR product was digested with PflMI and ClaI and purified by agarose gel electrophoresis in 3% NuSieve Agarose (FMC Corp.). Bacterial expression vector pSP26 (supra) was digested with PflMI and ClaI and the larger The PflMI-Clai PCR vector fragment was purified. fragment was ligated into the previously digested pSP26 containing the Amp gene. The ligation product was used to transform E. coli strain MC1061 and colonies were selected by ampicillin resistance. The resulting plasmid is denoted pSP26:Amp.

The F1 origin of replication from the mammalian expression vector pcDNAII (Invitrogen) was isolated in a 692 bp EarI fragment. Plasmid pcDNAII was digested with EarI and the resulting 692 bp fragment purified by agarose gel electrophoresis. EarI-NotI adapters were added to the 692 bp EarI fragment by ligation of two annealed oligonucleotide pairs, 179 + 180 and 181 + 182. The oligo pairs were annealed as described above.

179: GGCCGCTCTTCC

180: AAAGGAAGAGC

TAGAATTGC

"GCCGCAATT"

The oligonucleotide-ligated fragment was then ligated into the single NotI site of PSP26:Amp to yield the vector pSP26:Amp:F1.

B. Construction of vector pgIII

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The construction of pgIII is outlined in Figure 31. The portion of the phage geneIII protein gene contained by the PDW1 #14 phagemid vector was originally obtained as a PCR amplification product from vector m13mp8. A portion of m13mp8 geneIII encoding the carboxyl-terminal 158 amino acid residues of the geneIII product was isolated by PCR amplification of m13mp8 nucleotide residues 2307-2781 using PCR oligos 6162 and 6160.

6162: GCCGGATCCGCTATTTCCGGTGGCTCTGGTTCC

6160: GCCAAGCTTATTAAGACTCCTTATTACGCAG

The PCR oligos contain Bamil and HindIII restriction recognition sites such that PCR from m13mp8 plasmid DNA with the oligo pair yielded a 490 bp Bamil-HindIII fragment encoding the appropriate portion of geneIII. The PCR product was ligated between the Bamil and HindIII sites within the polylinker of PUC19 to yield plasmid pgIII.

C. Construction of pPhoA: KPI:gIII

Construction of pPhoA:KPI:gIII is outlined in Figure 32. A portion of the phoA signal sequence and KPI fusion encoded by the phage display vector PDW1 #14 originates with pPhoA:KPI:gIII. The 237 bp NdeI-HindIII fragment of pTW10:KPI encoding the entire phoA:KPI (1+57) fusion was isolated by preparative agarose gel electrophoresis, and inserted between the NdeI and HindIII sites of pUC19 to yield plasmid pPhoA:KPI. The 490 bp BamHI-HindIII fragment of pgIII encoding the C-terminal portion of the geneIII product was then isolated and ligated between the BamHI and HindIII sites of pPhoA:KPI to yield vector pPhoa:KPI:gIII. The pPhoA:KPI:gIII vector encodes a 236 amino acid residue

fusion of the phoA signal peptide, KPI (1→57) and the carboxyl-terminal portion of the geneIII product.

D. Construction of pLG1

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Construction of pLG1 is illustrated in Figure 33. The exact geneIII sequences contained in vector PDW1 #14 originate with phage display vector pLG1. A modified geneIII segment was generated by PCR amplification of the geneIII region from pgIII using PCR oligonucleotides 6308 and 6305.

6308: AGCTCCGATCTAGGATCCGGTGGTGGCTCTGGTTCCGGT 10

> GCAGCGGCCGTTAAGCTTATTAAGACTCCT 6305:

amplification from pgIII with these PCR oligonucleotides yielded a 481 bp BamHI-HindIII fragment encoding a geneIII product shortened by 3 amino acid residues at the amino-terminal portion of the segment of the geneIII fragment encoded by pgIII. A 161 bp NdeI-BamHI fragment was generated by PCR amplification from plasmid bacterial expression pTHW05 using oligonucleotides 6306 and 6307.

20 6306: GATCCTTGTGTCCATATGAAACAAAGC

> 6307: CACGTCGGTCGAGGATCCCTAACCACGGCCTTTAACCAG

The 161 bp NdeI-BamHI fragment and the 481 bp BamHI-HindIII fragment were gel-purified, and then ligated in a three-way ligation into PTW10 which had previously been digested with NdeI and HindIII. The resulting plasmid 25 pLG1 encodes a phoA signal peptide-insert-geneIII fusion for phage display purposes.

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E. Construction of pAL51

Construction of pAL51 is illustrated in Figure 34. Vector pAL51 contains the geneIII sequences of pLG1 which are to be incorporated in vector pDW1 #14.

A 1693 bp fragment of plasmid pBR322 was isolated, extending from the BamHI site at nucleotide 375 to the PvuII site at position 2064. Plasmid pLG1 was digested with Asp718I and BamHI, removing an 87 bp fragment. The overhanging Asp718I end was blunted by treatment with Klenow fragment, and the PvuII-BamHI fragment isolated from pBR322 was ligated into this vector, resulting in the insertion of a 1693 bp "stuffer" region between the Asp718I and BamHI sites. The 78 bp NdeI-Asp718I region of the resulting plasmid was removed and replaced with the annealed oligo pair 6512 + 6513.

6512: TATGAAACAAAGCACTATTGCACTGGCACTCTTACCGTTACTGTT
TACCCCGGTGACCAAAGCCCACGCTGAAG

6513: GTACCTTCAGCGTGGGCTTTGGTCACCGGGGTAAACAGTAACGGT
AAGAGTGCCAGTGCAATAGTGCTTTGTTTCA

The newly created 74 bp NdeI-Asp718I fragment encodes the phoA signal peptide, and contains a BstEII cloning site. The resulting plasmid is denoted pAL51.

F. Construction of pAL53

Construction of pAL53 is outlined in Figure 35. Plasmid pAL53 contributes most of the vector sequence of pDW1 #14, including the basic vector backbone with Amp gene, F1 origin, low copy number origin of replication, geneIII segment, phoA promotor and phoA signal sequence.

Plasmid pAL51 was digested with NdeI and HindIII and the resulting 2248 bp NdeI-HindIII fragment encoding the phoA signal peptide, stuffer region and geneIII region was isolated by preparative agarose gel electrophoresis. The NdeI-HindIII fragment was ligated into plasmid pSP26:Amp:Fl between the NdeI and HindIII sites, resulting in plasmid pAL52.

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The phoA promoter region and signal peptide was generated by amplification of a portion of the E. coli genome by PCR, using oligonucleotide primers 405 and 406.

405: CCGGACGCGTGGAGATTATCGTCACTG

5 406: GCTTTGGTCACCGGGGTAAACAGTAACGG

The resulting PCR product is a 332 bp MluI-BstEII fragment which contains the phoA promoter region and signal peptide sequence. This fragment was used to replace the 148 bp MluI-BstEII segment of PAL52, resulting in vector pAL53.

- G. Construction of pSP26:Amp:F1:PhoA:KPI:gIII

 Construction of pSP26:Amp:F1:PhoA:KPI:gIII is

 illustrated in Figure 36. This particular vector is the
 source of the KPI coding sequence found in vector pDW1

 #14. Plasmid pPhoa:KPI:gIII was digested with NdeI and
 HindIII, and the resulting 714 bp NdeI-HindIII fragment
 was purified, and then inserted into vector pSP26:Amp:F1
 between the NdeI and HindIII sites. The resulting
 plasmid is denoted pSP26:Amp:F1:PhoA:KPI:gIII.
- 20 H. Construction of pDW1 #14

 Construction of pDW1 #14 is illustrated in Figure 37.

 The sequences encoding KPI were amplified from plasmid pSP26:Amp:F1:PhoA:KPI:gIII by PCR, using oligonucleotide primers 424 and 425.
- 25 424: CTGTTTACCCCGGTGACCAAAGCCGAGGTGTGCTCTGAACAA
 425: AATAGCGGATCCGCACACTGCCATGCAGTACTCTTC

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The resulting 172 bp BstEII-BamHI fragment encodes most of KPI (1 \rightarrow 55). This fragment was used to replace the stuffer region in pAL53 between the BstEII and BamHI sites. The resulting plasmid, PDW1 #14, is the parent

To phage expraries the moding region (or the phoa-KP) (1-55)-geneIII fusion is shown in Figure 38.

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I. Construction of pDW1 14-2

Construction of pDWl 14-2 is illustrated in Figure 39. The first step in the construction of the KPI phage libraries in pDWl #14 was the replacement of the AgeI-BamHI fragment within the KPI coding sequence with a stuffer fragment. This greatly aids in preparation of randomized KPI libraries which are substantially free of contamination of phagemid genomes encoding wild-type KPI sequence.

Plasmid pDW1 #14 was digested with AgeI and BamHI, and the 135 bp AgeI-BamHI fragment encoding KPI was discarded. A stuffer fragment was created by PCR amplification of a portion of the PBR322 Tet gene, extending from the BamHI site at nucleotide 375 to nucleotide 1284, using oligo primers 266 and 252.

266: GCTTTAAACCGGTAGGTGGCCCGGCTCCATGCACC

252: CGAATTCACCGGTGTCATCCTCGGCACCGTCACCCT

The resulting 894 bp AgeI-BamHI stuffer fragment was then inserted into the AgeI/BamHI-digested pDW1 #14 to yield the phagemid vector pDW1 14-2. This vector was the starting point for construction of the randomized KPI libraries.

J. Construction of KPI Library 16-19

Construction of KPI Library 16-19 is outlined in Figure 40. Library 16-19 was constructed to display KPI-geneIII fusions in which amino acid positions Ala¹¹, Met¹³, Ile¹⁶ and Ser¹⁷ are randomized. For preparation of the library, plasmid pDW1 14-2 was digested with AgeI and BamHI to remove the stuffer region, and the resulting vector was purified by preparative agarose gel electrophoresis. Plasmid pDW1 #14 was used as template in a PCR amplification of the KPI region extending from the AgeI site to the BamHI site. The oligonucleotide primers used were 544 and 551.

544: GGGCTGAGACCGGTCCGTGCCGT (NNS) ACGCTGGTACTTTGACGTC

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551: GGAATAGCGGATCCGCACACTGCCATGCAG

Oligonucleotide primer 544 contains four randomized codons of the sequence NNS, where N represents equal mixtures of A/G/C/T and S an equal mixture of G or C. Each NNS codon thus encodes all 20 amino acids plus a single possible stop codon, in 32 different DNA sequences. PCR amplification from the wild-type KPI gene resulted in the production of a mixture of 135 bp AgeI-BamHI fragments all containing different sequences in the The PCR product was purified by randomized region. preparative agarose gel electrophoresis and ligated into the AgeI/BamHI digested PDW1 14-2 vector. The ligation mixture was used to transform E. coli Top10F' cells (Invitrogen) by electroporation according to manufacturer's directions. The resulting Library 16-19 contained approximately 400,000 independent clones. The potential size of the library, based upon the degeneracy of the priming PCR oligo #544 was 1,048,576 members. The expression unit encoded by the members of Library 16-19 is shown in Figure 41.

K. Selection of Library 16-19 with human plasma kallikrein

KPI phage were prepared and amplified by infecting transformed cells with M13K07 helper phage as described by Matthews et al., Science 260:1113 (1993). plasma kallikrein (Enzyme Research Laboratories, South Bend, IN), was coupled to Sepharose 6B resin. Prior to phage binding, the immobilized kallikrein resin was washed three times with 0.5 ml assay buffer (AB = 100mM Tris-HCl, pH 7.5, 0.5M NaCl, 5mM each of KCl, CaCl₂, 0.1% gelatin, and 0.05% Triton X-100). Approximately 5x10° phage particles of the amplified Library 16-19 in PBS, pH 7.5, containing 300mM NaCl and 0.1% gelatin, were bound to 50 µl kallikrein resin containing 15 pmoles of active human plasma kallikrein in a rotal volume of 250 µl. Phage were allowed to bind for

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in 0.5 ml AB. Bound phage were eluted sequentially by successive 5 minute washes: 0.5 ml 50mM sodium citrate, pH 6.0, 150mM NaCl; 0.5 ml 50mM sodium citrate, pH 4.0, 150mM NaCl; and 0.5 ml 50mM glycine, pH 2.0, 150mM NaCl. Eluted phage were neutralized immediately and phagemids from the pH 2.0 elution were titered and amplified for reselection. After three rounds of selection on kallikrein-Sepharose, phagemid DNA was isolated from 22 individual colonies and subjected to DNA sequence analysis.

The most frequently occurring randomized KPI region encoded: Ala^{15} - Ile^{16} - Phe^{17} . The phoA-KPI-geneIII region encoded by this class of selected KPI phage is shown in Figure 42. The KPI variant encoded by these phagemids is denoted KPI (1 \rightarrow 55; MISA, S17F).

L. Construction of pDD185 KPI (-4+57; M15A, S17F)
Figure 43 outlines the construction of pDD185 KPI
(-4+57; M15A, S17F). The sequences encoding KPI (1+55;
M15A, S17F) were moved from one phagemid vector, pDW1
20 . (16-19) 185, to the yeast expression vector so that the
KPI variant could be purified and tested.

Plasmid pTW113 encoding wild-type KPI (-4 \div 57) was digested with AgeI and BamHI and the 135 bp AgeI-BamHI fragment was discarded. The 135 bp AgeI-BamHI fragment of pDW1 (16-19) 185 was isolated and ligated into the yeast vector to yield plasmid pDD185, encoding α -factor fused to KPI (-4 \div 57; M15A, S17F). See Figure 44.

- M. Purification of KPI (-4+57; M15A, S17F) pDD185
 Transformation of yeast strain ABL115 with pDD185,
 induction of yeast cultures, and purification of KPI
 (-4+57; M15A, S17F) pDD185 was accomplished as described
 for the other KPI variants.
 - N. Construction of KPI Library 6 M15A, with residues 14, 16-18 random.
- Library 6 was constructed to display KPI-geneIII fusions in which amino acid positions Ala¹⁴, Ile¹⁶, Ser¹⁷

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and Arg¹⁸ are randomized, but position 15 was held constant as Ala. For preparation of the library, plasmid pDW1 #14 was used as template in a PCR amplification of the KPI region extending from the AgeI site to the BamHI site. The oligonucleotide primers used were 551 and 1003.

1003: GCTGAGACCGGTCCGTGCCGTNNSGCA(NNS),TGGTACTTTGACGTC

551: GGAATAGCGGATCCGCACACTGCCATGCAG

Oligonucleotide primer 1003 contained four randomized codons of the sequence NNS, where N represents equal mixtures of A/G/C/T and S an equal mixture of G or C. Each NNS codon thus encodes all 20 amino acids plus a single possible stop, in 32 different DNA sequences. PCR amplification from the wild-type KPI gene resulted in the production of a mixture of 135 bp Agel-BanHI fragments all containing different sequences in the randomized The PCR product was phenol extracted, ethanol precipitated, digested with BasHI and purified by preparative agarose gel electrophoresis. Plasmid pDW1 14-2 was digested with BamHI, phenol extracted and ethanol precipitated. The insert was ligated at high molar ratio to the vector which was then digested with AgeI to remove the stuffer region. The vector containing the insert was purified by agarose gel electrophoresis The resulting library contains and recircularized. approximately 5x10° independent clones.

O. Construction of KPI Library 7 — residues 14-18 random.

Library 7 was constructed to display KPI-geneIII fusions in which amino acid positions Ala¹⁶, Met¹⁵, Ile¹⁶, Ser¹⁷ and Arg¹⁸ are randomized. For preparation of the library, plasmid pDW1 #14 was used as template in a PCR

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GCTGAGACCGGTCCGTGCCGT(NNS),TGGTACTTTGACGTC 1179:

551: GGAATAGCGGATCCGCACACTGCCATGCAG

Oligonucleotide primer 1179 contains five randomized codons of the sequence NNS, where N represents equal mixtures of A/G/C/T and S an equal mixture of G or C. Each NNS codon thus encoded all 20 amino acids plus a single possible stop, in 32 different DNA sequences. PCR amplification from the wild-type KPI gene resulted in the production of a mixture of 135 bp AgeI-BamHI fragments all containing different sequences in the randomized region. The PCR product was phenol extracted, ethanol precipitated, digested with BamHI and purified by preparative agarose gel electrophoresis. Plasmid pDW1 14-2 was digested with BamHI, phenol extracted and ethanol precipitated. The insert was ligated at high molar ratio to the vector which was then digested with AgeI to remove the stuffer region. The vector containing the insert was purified by agarose gel electrophoresis The resulting library contains and recircularized. approximately 1x10⁷ independent clones.

Selection of Libraries 6 & 7 with human factor P.

KPI phage were prepared and amplified by infecting transformed cells with M13K07 helper phage (Matthews and Human factor XIIa (Enzyme Research Wells, 1993). Laboratories, South Bend, IN), was biotinylated as follows. Factor XIIa (0.5 mg) in 5mM sodium acetate pH 8.3 was incubated with Biotin Ester (Zymed) at room temperature for 1.5 h, then buffer-exchanged into assay buffer (AB). Approximately 1x1010 phage particles of each amplified Library 6 or 7 in PBS, pH 7.5, containing 300mM NaCl and 0.1% gelatin, were incubated with 50 pmoles of active biotinylated human factor XIIa in a total volume of 200 μ l. Phage were allowed to bind for 2 h at room temperature, with rocking. Following the binding period, 100 μ l Strepavidin Magnetic Particles (Boehringer

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Mannheim) were added to the mixture and incubated at room Separation of magnetic temperature for 30 minutes. particles from the supernatant and wash/elution buffers was carried out using MPC-E-1 Neodymium-iron-boron permanent magnets (Dynal). Unbound phage were removed by washing the magnetically bound biotinylated XIIa-phage complexes three times with 0.5 ml AB. Bound phage were eluted sequentially by successive 5 minute washes: 0.5 ml 50mM sodium citrate, pH 6.0, 150mM NaCl; 0.5 ml 50mM sodium citrate, pH 4.0, 150mM NaCl; and 0.5 ml 50mM glycine, pH 2.0, 150mM NaCl. Eluted phage were neutralized immediately and phagemids from the pH 2.0 elution were titered and amplified for reselection. After 3 or 4 rounds of selection with factor XIIa, phagemid DNA was isolated from individual colonies and subjected to DNA sequence analysis.

Sequences in the randomized regions were compared with one another to identify consensus sequences appearing more than once. From Library 6 a phagemid was identified which encoded MI5L, S17Y, R18H. From Library 7 a phagemid was identified which encoded MI5A, S17Y, R18H.

Q. Construction of pBG015 KPI (-4+57; M15L, S17Y, R18H), pBG022 (-4+57; M15A, S17Y, R18H)

The sequences encoding KPI (1-55; M15L, S17Y, R18H) and KPI (1-55; M17A, S17Y, R18H) were moved from the phagemid vectors to the yeast expression vector so that the KPI variant could be purified and tested.

Plasmid pTW113 encoding wild-type KPI (-4+57) was digested with AgeI and BamHI and the 135 bp AgeI-BamHI fragment was discarded. The 135 bp AgeI-BamHI fragment of the phagemid vectors were isolated and ligated into the yeast vector to yield plasmids pBG015 and pBG022, encoding alpha-factor fused to KPI (-4+57; M15L, S17Y, R18H), and KPI (-4+57; M15A, S17Y, R18H), respectively.

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R. Construction of pBG029 KPI (-4-57, T9V, M15L, S17Y, R18H)

Plasmid pBG015 was digested with XbaI and RsrII, and the larger of the two resulting fragments was isolated. An oligonucleotide pair (1593 + 1642) was phosphorylated, annealed and gel-purified as described previously.

1593: CTAGATAAAAGAGAGGTTGTTAGAGAGGTGTGCTCTGAACAAGCT GAGGTTG

1642: GACCAACCTCAGCTTGTTCAGAGCACACCTCTCTAA
CAACCTCTCTTTTAT

The annealed oligonucleotides were ligated into the XbaI and RsrII-digested pBG015, and the ligation product was used to transform E. coli strain MC1061 to ampicillin resistance. The resulting plasmid pBG029, encodes the 445 bp synthetic gene for the alpha-factor-KPI (-4÷57; T9V, M15L, S17F, R18H) fusion.

S. Construction of pBG033 KPI (-4→57; T9V, M15A, S17Y, R18H)

plasmid pBG022 was digested with XbaI and RsrII, and
the larger of the two resulting fragments was isolated.
An oligonucleotide pair (1593 + 1642) was phosphorylated,
annealed and gel-purified as described previously. The
annealed oligonucleotides were ligated into the XbaI and
RsrII-digested pBG022, and the ligation product was used
to transform E. coli strain MC1061 to ampicillin
resistance. The resulting plasmid pBG033, encodes the
445 bp synthetic gene for the alpha-factor-KPI (-4÷57;
T9V, M15A, S17F, R18H) fusion.

T. Selection of Library 16-19 with human factor Xa

KPI phage were prepared and amplified by infecting transformed cells with M13K07 helper phage (Matthews and Wells, 1993). Human factor Xa (Haematologic Technologies, Inc., Essex Junction, VT) was coupled to Sepharose 6B resin. Prior to phage binding, the

immobilized Xa resin was washed three times with 0.5 ml assay buffer (AB = 100mM Tris-HCl, pH 7.5, 0.5M NaCl, 5mM each of KCl, CaCl₂, MgCl₂, 0.1% gelatin, and 0.05% Triton Approximately 4x1010 phage particles of the amplified Library 16-19 in PBS, pH 7.5, containing 300mM NaCl and 0.1% gelatin, were bound to 50 μ l Xa resin in a total volume of 250 μ l. Phage were allowed to bind for 4 h at room temperature, with rocking. Unbound phage were removed by washing the Xa resin three times in 0.5 Bound phage were eluted sequentially by successive 5 minute washes: 0.5 ml 50mM sodium citrate, pH 6.0, 150mM NaCl; 0.5 ml 50mM sodium citrate, pH 4.0 150mM NaCl; and 0.5 ml 50mM glycine, pH 2.0, 150mM NaCl. Eluted phage were neutralized immediately and phagemids from the pH 2.0 elution were titered and amplified for After three rounds of selection on Xareselection. Sepharose, phagemid DNA was isolated and subjected to DNA sequence analysis.

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Sequences in the randomized Ala^{14} -Ser¹⁷ region were compared with one another to identify consensus sequences appearing more than once. A phagemid was identified which encoded KPI (1 \Rightarrow 55; M15L, I16F, S17K).

U. Construction of pDD131 KPI (-4→57; M15L, I16F, S17K)

The sequences encoding KPI (1-55; M15L, I16F, S17K) were moved from the phagemid vector to the yeast expression vector so that the KPI variant could be purified and tested.

Plasmid pTW113 encoding wild-type KPI (-4-57) was digested with AgeI and BamHI and the 135 bp AgeI-BamHI fragment was discarded. The 135 bp AgeI-BamHI fragment of the phagemid vector was isolated and ligated into the yeast vector to yield plasmid pDD131, encoding alphafactor fused to KPI (-4-57; M15L, I16F, S17K).

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V. Construction of pDD134 KPI (-4→57; M15L, I16F, S17K, G37Y)

plasmid pDD131 was digested with AatI and BamHI, and the larger of the two resulting fragments was isolated. An oligonucleotide pair (738 + 739) was phosphorylated, annealed and gel-purified as described previously.

- 738: CACTGAAGGTAAGTGCGCTCCATTCTTTTACGGCGGTTGCTACGGCAACCGT
 AACAACTTTGACACTGAAGAGTACTGCATGGCAGTGTGCG
- 739: GATCCGCACACTGCCATGCAGTACTCTTCAGTGTCAAAGTTGTTACGGTTGC CGTAGCAACCGCCGTAAAAGAATGGAGCGCACTTACCTTCAGTGACGT

The annealed oligonucleotides were ligated into the AatI and BamHI-digested pDD131, and the ligation product was used to transform E. coli strain MC1061 to ampicillin resistance. The resulting plasmid pDD134, encodes the 445 bp synthetic gene for the alpha-factor-KPI (-4 \rightarrow 57; M15L, I16F, S17K, G37Y) fusion.

W. Construction of pDD135 KPI (-4-57; M15L, I16F, S17K, G37L)

Plasmid pDD131 was digested with AatII and BamHI, and the larger of the two resulting fragments was isolated. An oligonucleotide pair (724 + 725) was phosphorylated, annealed and gel-purified as described previously.

- 738: CACTGAAGGTAAGTGCGCTCCATTCTTTTACGGCGGTTGCTACGGCAACCGT
 AACAACTTTGACACTGAAGAGTACTGCATGGCAGTGTGCG
- 25 739: GATCCGCACACTGCCATGCAGTACTCTTCAGTGTCAAAGTTGTTACGGTTGC
 CGTAGCAACCGCCGTAAAAGAATGGAGCGCACTTACCTTCAGTGACGT

The annealed oligonucleotides were ligated into the AatII and BamHI-digested pDD131, and the ligation product was used to transform E. coli strain MC1061 to ampicillin resistance. The resulting plasmid pDD135, encodes the

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445 bp synthetic gene for the alpha-factor-KPI (-4 \Rightarrow 57; M15L, I16F, S17K, G37L) fusion.

Example 4. Kinetic analysis of KPI(-4+57) variants

The concentrations of active human plasma kallikrein, factor XIIa, and trypsin were determined by titration with p-nitrophenyl p'-guanidinobenzoate as described by Bender et al., supra, and Chase et al., Biochem. Biophys. Res. Commun. 29:508 (1967). Accurate concentrations of active KPI(-4→57) inhibitors were determined by titration of the activity of a known amount of active-site-titrated For testing against kallikrein and trypsin, each KPI(-4-57) variant (0.5 to 100nM) was incubated with protease in low-binding 96-well microtiter plates at 30°C for 15-25 min, in 100mM Tris-HCl, pH 7.5, with 500mM NaCl, 5mM KCl, 5mM CaCl2, 5mM MgCl2, 0.1% Difco gelatin, and 0.05% Triton X-100. Chromogenic synthetic substrate was then be added, and initial rates at 30°C recorded by the SOFTmax kinetics program via a THERMOmax microplate reader (Molecular Devices Corp., Menlo Park, CA). substrates used were N-a-benzoyl-L-Arg p-nitroanilide (1mM) for trypsin (20nM), and N-benzoyl-Pro-Phe-Arg pnitroanilide (0.3mM) for plasma kallikrein (1mM). Enzfitter (Elsevier) program was used both to plot fractional activity (i.e., activity with inhibitor, divided by activity without inhibitor), a, versus total concentration of inhibitor, I, and to calculate the dissociation constant of the inhibitor (K) by fitting the curve to the following equation:

$$a=1-\frac{[E]_{c}+[I]_{c}+K_{i}-\sqrt{([E]_{c}+[I]_{c}+K_{i})^{2}-4[E]_{c}[I]_{c}}}{2[E]_{c}}$$

nown foure the fost fotent fariants To -4-57; M15A, S17F) DD185 and KPI -4-57; M15A, S17Y; TW6166 are 115-fold and 100-fold more potent,

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respectively, as a human kallikrein inhibitor than wild-type KPI $(-4 \rightarrow 57)$. The least potent variant, KPI $(-4 \rightarrow 57)$; I16H, S17W) TW6185 is still 35-fold more potent than wild-type KPI.

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For testing against factor XIIa, essentially the same reaction conditions were used, except that the substrate was N-benzoyl-Ile-Glu-Gly-Arg p-nitroaniline hydrochloride and its methyl ester (obtained from Pharmacia Hepar, Franklin, OH), and corn trypsin inhibitor (Enzyme Research Laboratories, South Bend, IN) was used as a control inhibitor. Factor XIIa was also obtained from Enzyme Research Laboratories.

Various data for inhibition of the serine proteases of interest kallikrein, plasmin, and factors Xa, XIa, and XIIa by a series of KPI variants are given in Figure 46. The results indicate that KPI variants can be produced that can bind to and preferably inhibit the activity of serine proteases. The results also indicate that the peptides of the invention may exhibit the preferable more potent and specific inhibition of one or more serine proteases of interest.

Example 5. Effect of KPI variant KPI185-1 on postoperative bleeding

A randomized, double-blinded study using an acute porcine cardiopulmonary bypass (CPB) model was used to investigate the effect of KPI185-1 on postoperative bleeding. Sixteen pigs (55-65 kg) underwent 60 minutes of hypothermic (28°C) open-chest CPB with 30 minutes of cardioplegic cardiac arrest. Pigs were randomized against a control solution of physiological saline (NS; n=8) or KPI-185 (n=8) groups. During aortic crossclamping, the tricuspid valve was inspected through an atriotomy which was subsequently repaired. Following reversal of heparin with protamine, dilateral thoracostomy tubes were placed and shed blood collected for 3 hours. Shed blood volume and hemoglobin (Hgb) loss were calculated from total chest tube output and residual intrathoracic blood at time of sacrifice.

Total blood loss was significantly reduced in the KPI185-1 group (245.75 \pm 66.24 ml vs. 344.25 \pm 63.97 ml, p=0.009). In addition, there was a marked reduction in total Hgb loss in the treatment group (13.59 \pm 4.26 gm vs. 23.61 \pm 4.69 gm, p=0.0005). Thoracostomy drainage Hgb was significantly increased at 30 and 60 minutes in the control group [6.89 \pm 1.44 vs. 4.41 \pm 1.45 gm/dl (p=0.004) and 7.6 \pm 1.03 vs. 5.26 \pm 1.04 gm/dl (p=0.0002), respectively]. Preoperative and post-CPB hematocrits were not statistically different between the groups. These results are shown in graphical form in Figures 47-50.

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The invention has been disclosed broadly and illustrated in reference to representative embodiments described above. Those skilled in the art will recognize that various modifications can be made to the present invention without departing from the spirit and scope thereof.

What Is Claimed Is:

1. A protease inhibitor comprising the sequence:

X¹-Val-Cys-Ser-Glu-Gln-Ala-Glu-X¹-Gly-X³
Cys-Arg-Ala-X⁴-X⁵-X⁶-X⁷-Trp-Tyr-Phe-Asp
Val-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-X⁶
Tyr-Gly-Gly-Cys-X⁶-X¹⁰-X¹¹-X¹²-Asn-Asn-Phe
Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-Val-Cys
Gly-Ser-Ala-Ile,

wherein:

 \mathbf{X}^{i} is selected from Glu-Val-Val-Arg-Glu-, Asp, or Glu:

 X^2 is selected from Thr, Val, Ile and Ser;

X3 is selected from Pro and Ala;

 X^4 is selected from Arg, Ala, Leu, Gly, or Met;

X' is selected from Ile, His, Leu, Lys, Ala, or Phe;

X' is selected from Ser, Ile, Pro, Phe, Tyr, Trp, Asn, Leu, His, Lys, or Glu;

 X^7 is selected from Arg, His, or Ala;

X4 is selected from Phe, Val, Leu, or Gly;

X* is selected from Gly, Ala, Lys, Pro, Arg, Leu,
Met, or Tyr;

X10 is selected from Ala, Arg, or Gly;

Xⁱⁱ is selected from Lys, Ala, or Asn;

X12 is selected from Ser, Ala, or Arg;

provided that:

when X' is Arg, X' is Ile;

when X^0 is Arg, X^4 is Ala or Leu; when X^0 is Tyr, X^4 is Ala or X^3 is His; and

either X^5 is not Ile; or X^6 is not Ser; or X^6 is not Leu, Phe, Met, Tyr, or Asn; or X^{10} is not Gly; or X^{11} is not Asn; or X^{12} is not Arg.

2. A protease inhibitor comprising the sequence: X¹-Val-Cys-Ser-Glu-Gln-Ala-Glu-Thr-Gly-Pro-Cys-Arg-Ala-X²-X³-X⁴-Arg-Trp-Tyr-Phe-Asp-Val-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-Phe-Tyr-Gly-Gly-Cys-X⁵-Gly-Asn-Arg-AsnAsn-Phe-Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-Val-Cys-Gly-Ser-Ala-Ile,

wherein:

 \mathbf{X}^{t} is selected from Glu-Val-Val-Arg-Glu-, Asp, or Glu;

X2 is selected from Ala, Leu, Gly, or Met;

X3 is selected from Ile, His, Leu, Lys, Ala, or Phe;

X⁴ is selected from Ser, Ile, Pro, Phe, Tyr, Trp,
Asn, Leu, His, Lys, or Glu;

 $\mathbf{X}^{\mathbf{J}}$ is selected from Gly, Ala, Lys, Pro, Arg, Leu, Met, or Tyr;

provided that:

when X^5 is Arg, X^2 is Ala or Leu; when X^5 is Tyr, X^2 is Ala or X^3 is His; and

either X^3 is not Ile; or X^4 is not Ser; or X^5 is not Leu, Phe, Met, Tyr, or Asn.

3. A protease inhibitor comprising the sequence:

Glu-Val-Arg-Glu-Val-Cys-Ser-Glu-Gln-Ala-Glu-Thr-Gly-Pro-Cys-Arg-Ala-X¹-X²-X³-Arg-Trp-Tyr-Phe-Asp-Val-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-Phe-Tyr-Gly-Gly-Cys-X⁴-Gly-Asn-Arg-Asn-Asn-Phe-Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-Val-Cys-Gly-Ser-Ala-Ile,

wherein:

X' is selected from Ala, Leu, Gly, or Met;

X² is selected from Ile, His, Leu, Lys, Ala, or Phe;

X³ is selected from Ser, Ile, Pro, Phe, Tyr, Trp, Asn, Leu, His, Lys, or Glu;

X is selected from Gly, Arg, Leu, Met, or Tyr; provided that:

when X1 is Ala, X2 is Ile, His, or Leu;

when X' is Leu, X2 is Ile or His;

when X^i is Leu and X^2 is Ile, X^3 is not Ser;

when X' is Gly, X2 is Ile;

when X' is Arg X' is Ala or Leu-

ner.

Ser, or X" is not Gly.

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- 4. A protease inhibitor according to claim 1, wherein at least two amino acid residues selected from the group consisting of X^4 , X^5 , X^6 , and X^7 differ from the residues found in the naturally occurring sequence of KPI.
- 5. A protease inhibitor according to claim 1, wherein X^1 is Asp or Glu, X^2 is Thr, X^3 is Pro, and X^{12} is Ser.
- 6. A protease inhibitor according to claim 5, wherein X^1 is Glu, X^2 is Thr, X^3 is Pro, X^4 is Met, X^5 is Ile, X^6 is Ser, X^7 is Arg, x^8 is Phe, X^9 is Gly, X^{10} is Gly, and X^{11} is Asn.
- 7. A protease inhibitor according to claim 5, wherein X^1 is Asp, X^2 is Thr, X^3 is Pro, X^4 is Arg, X^5 is Ile, X^6 is Ile, X^7 is Arg, x^8 is Val, X^9 is Arg, X^{10} is Ala, and X^{11} is Lys.
- 8. A protease inhibitor according to claim 1, wherein X^1 is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Pro, X^4 is Met, X^5 is Ile, X^6 is Ser, X^7 is Arg, x^3 is Phe, X^9 is Gly, X^{10} is Gly, X^{11} is Asn, and X^{12} is Ala.
- 9. A protease inhibitor according to claim 1, wherein X^i is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Pro, X^4 is Met, X^5 is Ile, X^6 is Ser, X^7 is Arg, x^5 is Phe, X^9 is Gly, X^{10} is Gly, X^{11} is Ala, and X^{12} is Arg.
- 10. A protease inhibitor according to claim 1, wherein X^1 is Glu, X^2 is Thr, X^3 is Pro, X^4 is Met, X^5 is Ile, X^4 is Ser, X^7 is Arg, x^8 is Phe, X^9 is Gly, X^{10} is Ala, X^{11} is Asn, and X^{12} is Arg.
- 11. A protease inhibitor according to claim 1, wherein X^i is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Pro, X^4 is Met, X^3 is Ile, X^6 is Ser, X^7 is Arg, X^3 is Phe, X^6 is Gly, X^{10} is Arg, X^{11} is Asn, and X^{12} is Arg.

- 12. A protease inhibitor according to claim 1, wherein X^1 is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Pro, X^4 is Met, X^5 is Ile, X^6 is Ser, X^7 is Arg, x^8 is Val, Leu, or Gly, X^9 is Gly, X^{10} is Gly, X^{11} is Asn, and X^{12} is Arg.
- 13. A protease inhibitor according to claim 1, wherein X^i is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Pro, X^4 is Met, X^3 is Ile, X^4 is Ser, X^7 is Ala, x^3 is Phe, X^6 is Gly, X^{10} is Gly, X^{11} is Asn, and X^{12} is Arg.
- 14. A protease inhibitor according to claim 1, wherein X^1 is Glu-Val-Val-Arg-Glu-, X^2 is Thr, Val, or Ser, X^3 is Pro, X^4 is Ala or Leu, X^5 is Ile, X^6 is Tyr, X^7 His, X^8 is Phe, X^9 is Gly, X^{10} is Gly, X^{11} is Ala, and X^{12} is Arg.
- 15. A protease inhibitor according to claim 14, wherein X^2 is Thr, and X^4 is Ala.
- 16. A protease inhibitor according to claim 14, wherein X^2 is Thr, and X^4 is Leu.
- 17. A protease inhibitor according to claim 14, wherein X^2 is Val, and X^4 is Ala.
- 18. A procease inhibitor according to claim 14, wherein X^2 is Ser, and X^4 is Ala.
- 19. A protease inhibitor according to claim 14, wherein X^2 is Val, and X^4 is Leu.
- 20. A protease inhibitor according to claim 14, wherein X^2 is Ser, and X^4 is Leu.
- 21. A protease inhibitor according to claim 1, wherein X^1 is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Pro,

- 22. A protease inhibitor according to claim 1, wherein X^i is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Pro, X^4 is Leu, X^3 is Phe, X^6 is Lys, X^7 is Arg, X^6 is Phe, X^6 is Tyr, X^{10} is Gly, X^{11} is Ala, and X^{12} is Arg.
- 23. A protease inhibitor according to claim 1, wherein X^1 is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Pro, X^4 is Leu, X^5 is Phe, X^6 is Lys, X^7 is Phe, X^6 is Phe, Phe,
- 24. A protease inhibitor according to claim 2, wherein X^i is Glu, X^2 is Met, X^3 is Ile, X^4 is Ile, and X^5 is Gly.
- 25. A protease inhibitor according to claim 3, wherein X^i is Met, X^3 is Ser, and X^4 is Gly.
- 26. A protease inhibitor according to claim 25, wherein X^2 is selected from His, Ala, Phe, Lys, and Leu.
- 27. A protease inhibitor according to claim 26, wherein X^2 is His.
- 28. A protease inhibitor according to claim 27, wherein X^2 is Ala.
- 29. A protease inhibitor according to claim 27, wherein X^2 is Phe.
- 30. A protease inhibitor according to claim 27, wherein X^2 is Lys.
- 31. A protease inhibitor according to claim 27, wherein X^2 is Leu.
- 32. A protease inhibitor according to claim 3, wherein X^i is Met, X^2 is Ile, and X^i is Gly.

- 33. A protease inhibitor according to claim 32, wherein X^3 is Ile.
- 34. A protease inhibitor according to claim 32, wherein X' is Pro.
- 35. A protease inhibitor according to claim 32, wherein X^3 is Phe.
- 36. A protease inhibitor according to claim 32, wherein X^3 is Tyr.
- 37. A protease inhibitor according to claim 32, wherein X^3 is Trp.
- 38. A protease inhibitor according to claim 32, wherein X^3 is Asn.
- 39. A protease inhibitor according to claim 32, wherein X^3 is Leu.
- 40. A protease inhibitor according to claim 32, wherein X^3 is Lys.
- 41. A protease inhibitor according to claim 32, wherein X^3 is His.
- 42. A protease inhibitor according to claim 32, wherein X^3 is Glu.
- 43. A protease inhibitor according to claim 3, wherein \mathbf{X}^i is Ala.
- 44. A protease inhibitor according to claim 43, wherein X^2 is Ile.

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- 46. A protease inhibitor according to claim 44, wherein X^3 is Tyr, and X^4 is Gly.
- 47. A protease inhibitor according to claim 44, wherein X^3 is Trp, and X^4 is Gly.
- 48. A protease inhibitor according to claim 44, wherein X^3 is Ser or Phe, and X^4 is Arg or Tyr.
- 49. A protease inhibitor according to claim 43, wherein X^2 is His or Leu, X^3 is Phe, and X^4 is Gly.
- 50. A protease inhibitor according to claim 3, wherein X^{l} is Leu.
- 51. A protease inhibitor according to claim 50, wherein X^2 is His, X^3 is Asn or Phe, and X^4 is Gly.
- 52. A protease inhibitor according to claim 50, wherein X^2 is Ile, X^3 is Pro, and X^4 is Gly.
- 53. A protease inhibitor according to claim 3, wherein X^i is Gly, X^2 is Ile, X^3 is Tyr, and X^4 is Gly.
- 54. A protease inhibitor according to claim 3, wherein X^i is Met, X^2 is His, X^3 is Ser, and X^4 is Tyr.
- 55. An isolated DNA molecule comprising a DNA sequence encoding a protease inhibitor according to claim 1.
- 56. An isolated DNA molecule according to claim 55, operably linked to a regulatory sequence that controls expression of the coding sequence in a host cell.
- 57. An isolated DNA molecule according to claim 56. further comprising a DNA sequence encoding a secretory signal peptide.

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A See III

- 58. An isolated DNA molecule according to claim 57, wherein said secretory signal peptide comprises the signal sequence of yeast alpha-mating factor.
- 59. A host cell transformed with a DNA molecule according to claim 55.
- 60. A host cell according to claim 59, wherein said host cell is *E. coli* or a yeast cell.
- 61. A host cell according to claim 60, wherein said yeast cell is selected from Pichia pastoris and Saccharomyces cerevisiae.
- 62. A method for producing a protease inhibitor, comprising the steps of culturing a host cell according to claim 59 and isolating and purifying said protease inhibitor.
- 63. A pharmaceutical composition, comprising a protease inhibitor according to claim 1, together with a pharmaceutically acceptable sterile vehicle.
- 64. A method of treatment of a clinical condition associated with increased activity of one or more serine proteases, comprising administering to a patient suffering from said clinical condition an effective amount of a pharmaceutical composition according to claim 63.
- 65. The method of treatment of claim 64, wherein said clinical condition is blood loss during surgery.
- 66. A method for inhibiting the activity of serine proteases of interest in a mammal comprising administering a therapeutically effective dose of a

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- 67. The method of claim 66, wherein said serine proteases are selected from the group consisting of: kallikrein; chymotrypsins A and B; trypsin; elastase; subtilisin; coagulants and procoagulants, particularly those in active form, including coagulation factors such as factors VIIa, IXa, Xa, XIa, and XIIa; plasmin; thrombin; proteinase-3; enterokinase; acrosin; cathepsin; urokinase; and tissue plasminogen activator.
 - 68. A protease inhibitor comprising the sequence:

 Xi-Val-Cys-Ser-Glu-Gln-Ala-Glu-Xi-Gly-ProCys-Arg-Ala-Xi-Xi-Xi-Xi-Trp-Tyr-Phe-AspVal-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-PheTyr-Gly-Gly-Cys-Xi-Gly-Asn-Arg-Asn-AsnPhe-Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-ValCys-Gly-Ser-Ala-Ile,

wherein:

X¹ is selected from Glu-Val-Arg-Glu-, Asp, or Glu;

X2 is selected from Thr, Val, Ile and Ser;

X3 is selected from Arg, Ala, Leu, Gly, or Met;

X4 is selected from Ile, His, Leu, Lys, Ala, or Phe;

X³ is selected from Ser, Ile, Pro, Phe, Tyr, Trp,
Asn, Leu, His, Lys, or Glu;

X4 is selected from Arg, His, or Ala; and

 \mathbf{X}^{7} is selected from Gly, Ala, Lys, Pro, Arg, Leu, Met, or Tyr.

- 69. A protease inhibitor according to claim 68, wherein at least two amino acid residues selected from the group consisting of X^3 , X^4 , X^5 , and X^6 differ from the residues found in the naturally occurring sequence of KPI.
- 70. A protease inhibitor according to claim 68, wherein X^1 is Glu-Val-Val-Arg-Glu-, X^2 is Thr, Val, or Ser, X^3 is Ala or Leu, X^4 is Ile, X^5 is Tyr, X^6 is His and X^7 is Gly.

- 71. A protease inhibitor according to claim 70, wherein X^2 is Thr, and X^3 is Ala.
- 72. A protease inhibitor according to claim 70, wherein X^2 is Thr, and X^3 is Leu.
- 73. A protease inhibitor according to claim 70, wherein X^2 is Val, and X^3 is Ala.
- 74. A protease inhibitor according to claim 70, wherein X^2 is Ser, and X^3 is Ala.
- 75. A protease inhibitor according to claim 70, wherein X^2 is Val, and X^3 is Leu.
- 76. A protease inhibitor according to claim 70, wherein X^2 is Ser, and X^3 is Leu.
- 77. A protease inhibitor according to claim 68, wherein X^1 is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Leu, X^4 is Phe, X^3 is Lys, X^4 is Arg and X^7 is Gly.
- 78. A protease inhibitor according to claim 68, wherein X^1 is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Leu, X^4 is Phe, X^5 is Lys, X^4 is Arg and X^7 is Tyr.
- 79. A protease inhibitor according to claim 68, wherein X^1 is Glu-Val-Val-Arg-Glu-, X^2 is Thr, X^3 is Leu, X^4 is Phe, X^5 is Lys, X^4 is Arg and X^7 is Leu.
 - 80. A protease inhibitor comprising the sequence:

 X¹-Val-Cys-Ser-Glu-Gln-Ala-Glu-Thr-GlyPro-Cys-X²-Ala-X³-X⁴-X⁵-X⁶-Trp-Tyr-Phe-AspVal-Thr-Glu-Gly-Lys-Cys-Ala-Pro-Phe-PheTyr-Gly-Gly-Cys-Gly-Gly-Asn-Arg-Asn-AsnPhe-Asp-Thr-Glu-Glu-Tyr-Cys-Met-Ala-Val-

X1 is selected from Glu-Val-Val-Arg-Glu- and Asp-Val-Val-Arg-Glu-;

 X^2 is selected from Arg and Lys;

X' is selected from Met, Arg, Ala, Leu, Ser, Val;

X' is selected from Ile and Ala;

X' is selected from Ser, Ile, Ala, Pro, Phe, Tyr, and Trp; and

X' is selected from Arg, Ala, His, Gln, and Thr; provided that:

when X¹ is Arg, X³ is Leu, and X⁴ is Ile, X⁵ cannot be Ser; and also provided that either X' is not Met; or X' is not Ile; or X' is not Ser; or X' is not Arg.

- 81. A protease inhibitor according to claim 80, wherein X' is selected from Phe, Tyr and Trp.
- 82. A protease inhibitor according to claim 80, wherein X' is Ile.
- 83. A protease inhibitor according to claim 82, wherein X2 is Lys.
- 84. A protease inhibitor according to claim 83, wherein X3 is Met.
- 85. A protease inhibitor according to claim 84, wherein X' is Ser.
- 86. A protease inhibitor according to claim 84, wherein X' is Ile.
- 87. A protease inhibitor according to claim 83, wherein X3 is Arg.
- 88. A protease inhibitor according to claim 87, wherein X' is Ser.
- 89. A protease inhibitor according to claim 87, wherein X' is Ile.

90. A procease inhibitor according to claim 82, wherein \mathbf{X}^2 is Arg.

- 91. A protease inhibitor according to claim 90, wherein X^3 is Arg or Met, and X^5 is Ser or Ile.
- 92. A protease inhibitor according to claim 91, wherein X^3 is Arg.
- 93. A protease inhibitor according to claim 92, wherein $\mathbf{X}^{\mathbf{J}}$ is Ser.
- 94. A protease inhibitor according to claim 92, wherein X^3 is Ile.
- 95. A protease inhibitor according to claim 91, wherein X^3 is Met.
- 96. A protease inhibitor according to claim 95, wherein $\mathbf{X}^{\mathbf{5}}$ is Ser.
- 97. A protease inhibitor according to claim 95, wherein X⁵ is Ile.
- 98. A protease inhibitor according to claim 82, wherein X^3 is Ala.
- 99. A protease inhibitor according to claim 82, wherein X³ is Leu.

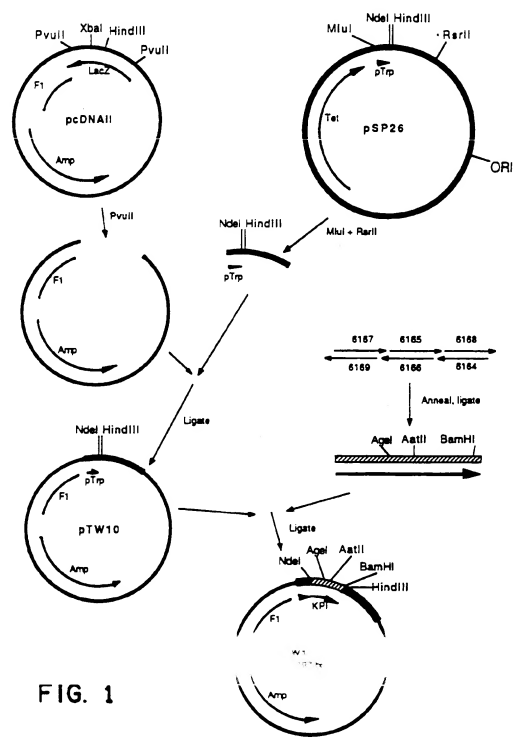
- 103. A protease inhibitor according to claim 82, wherein X^3 is Phe.
- 104. A protease inhibitor according to claim 82, wherein \mathbf{X}^{s} is Tyr.
- 105. A protease inhibitor according to claim 82, wherein X^3 is Trp.
- 106. A protease inhibitor according to claim 104, wherein X^3 is Ala or Leu.
- 107. A protease inhibitor according to claim 106, wherein X^3 is Ala.
- 108. A protease inhibitor according to claim 106, wherein X^3 is Leu.
- 109. A protease inhibitor according to claim 105, wherein X^3 is Ala.
- 110. A protease inhibitor according to claim 109, wherein X^5 is His.
- 111. A protease inhibitor according to claim 109, wherein X^5 is Gln.
- 112. A protease inhibitor according to claim 109, wherein X⁵ is Thr.
- 113. An isolated DNA molecule comprising a DNA sequence encoding a protease inhibitor according to claim 80.
- 114. An isolated DNA molecule according to claim 113, operably linked to a regulatory sequence that controls expression of the coding sequence in a host cell.

115. An isolated DNA molecule according to claim 114, further comprising a DNA sequence encoding a secretory signal peptide.

- 116. An isolated DNA molecule according to claim 115, wherein said secretory signal peptide comprises the signal sequence of yeast alpha-mating factor.
- 117. A host cell transformed with a DNA molecule according to claim 113.
- 118. A host cell according to claim 117, wherein said host cell is E. coli or a yeast cell.
- 119. A host cell according to claim 118, wherein said yeast cell is selected from Pichia pastoris and Saccharomyces cerevisiae.
- 120. A method for producing a protease inhibitor, comprising the steps of culturing a host cell according to claim 117 and isolating and purifying said protease inhibitor.
- 121. A pharmaceutical composition, comprising a protease inhibitor according to claim 80, together with a pharmaceutically acceptable sterile vehicle.
- 122. A method of treatment of a clinical condition associated with increased activity of one or more serine proteases, comprising administering to a patient suffering from said clinical condition an effective amount of a pharmaceutical composition according to claim 121.
 - 123. The method of treatment of claim 122, wherein
- 124. A method for inhibiting the activity of serine proteases of interest in a mammal comprising

administering a therapeutically effective dose of a pharmaceutical composition according to claim 121.

- 125. The method of claim 124, wherein said serine proteases are selected from the group consisting of: kallikrein; chymotrypsins A and B; trypsin; elastase; subtilisin; coagulants and procoagulants, particularly those in active form, including coagulation factors such as factors VIIa, IXa, Xa, XIa, and XIIa; plasmin; thrombin; proteinase-3; enterokinase; acrosin; cathepsin; urokinase; and tissue plasminogen activator.
- 126. A protease inhibitor according to claim 81, wherein X^4 is Ile.
- 127. A protease inhibitor according to claim 126, wherein X^5 is Phe.
- 128. A protease inhibitor according to claim 126, wherein \mathbf{X}^{S} is Tyr.
- 129. A protease inhibitor according to claim 126, wherein \mathbf{X}^{5} is Trp.
- 130. A protease inhibitor according to claim 128, wherein X^3 is Ala or Leu.
- 131. A protease inhibitor according to claim 130, wherein X^3 is Ala.
- 132. A protease inhibitor according to claim 130, wherein X^3 is Leu.
- 133. A protease inhibitor according to claim 129, wherein X^3 is Ala.

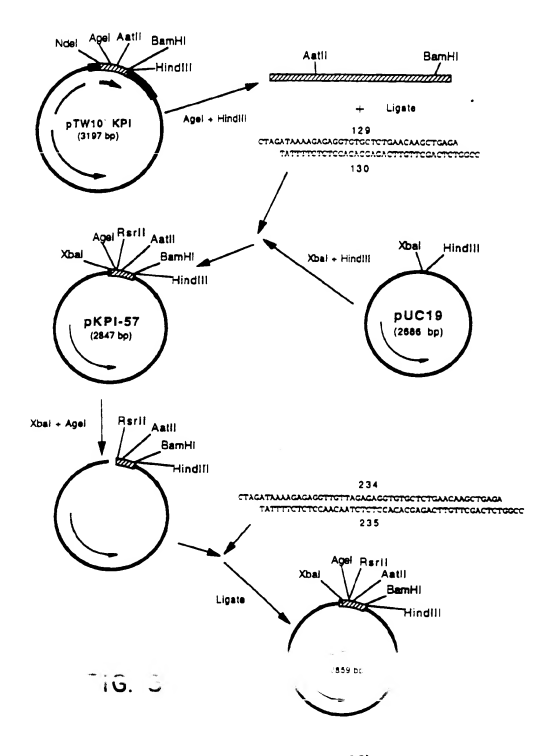


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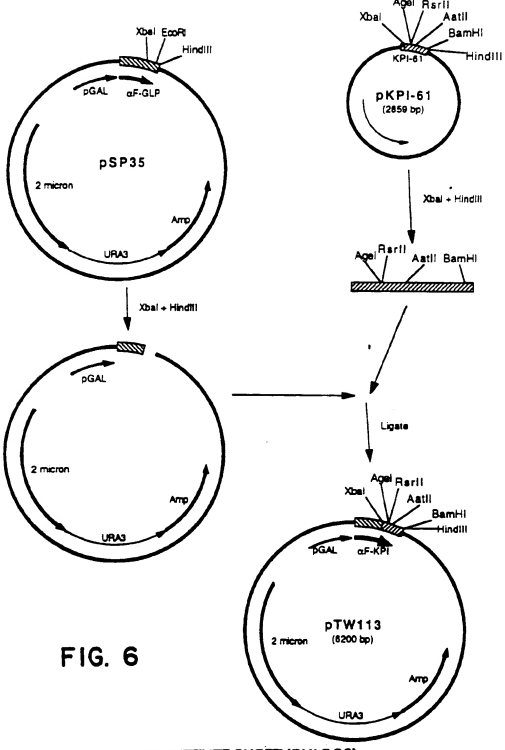
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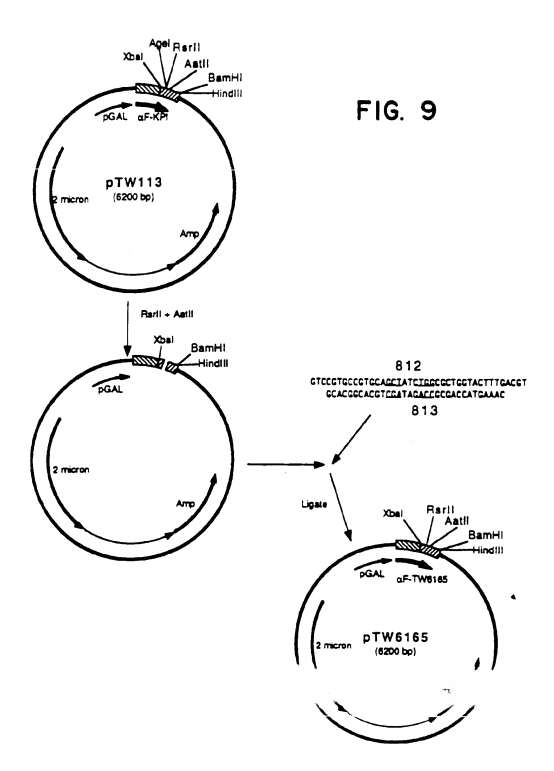
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FIG. 7

α-factor ATG AGA TIT CCT TCA ATT TTT ACT GCA GTT TTA TTC GCA GCA TCC TCC GCA TTA GCT TAC TOT AMA GGA AGT TAM AMA TGA COT CAM MAT MAG COT COT AGG AGG COT MAT COM Met Arg Phe Pro Ser lie Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG PAIR Pro Val Asn Thr Thr Thr Glu Asp Glu Thr Air Gin Ile Pro Air Glu Air Val ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG > lie Gly Tyr Leu Asp Leu Glu Gly Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn AGC AGA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT ► Ser Thr Asn Asn Gly Leu Leu Phe lie Asn Thr Thr !le Ala Ser lie Ala Ala Lys KPI(-4-57) GAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA CTT CTT CCC CAT AGA GAT CTA TTT TCT CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT FGIU GIU GIV Val Ser Leu Asp Lys Arg GIU Val Val Arg GIU Val Cys Ser GIU GIN Rsrll Agel Aatll GCT GAG ACC GGT CCG TGC CGT GCA ATG ATC TCC CGC TGG TAC TTT GAC GTC ACT GAA CGA CTC TGG CCA GGC ACG GCA CGT TAC TAG AGG GCG ACC ATG AAA CTG CAG TGA CTT PAIR GIU Thr Gly Pro Cys Arg Ala Met IIe Ser Arg Trp Tyr Phe Asp Val Thr Glu GST AAG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC AAC CGT AAC AAC TTT GAC CCA TTC ACG CGA GGT AAG AAA ATG CCG CCA ACG CCG CCG TTG GCA TTG TTG AAA CTG FGIY LYS CYS ALE Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp BamHi HindIII ACT GAA GAG TAC TGC ATG GCA GTG TGC GGA TCC GCT ATT TAA GCT T TGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A Thr Glu Glu Tyr Cys Met Ala Val Cys Gly Ser Ala ile

FIG. 8

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pTW 6165

FIG. 10

α-factor ATG AGA TIT OCT TOA ATT TIT ACT GOA GIT TIA TITC GOA GOA TOO TOO GOA TIA GOT TAC TOT AAA GGA AGT TAA AAA TGA CGT CAA AAT AAG CGT CGT AGG AGG CGT AAT CGA Met Arg Phe Pro Ser IIe Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG ▶Ala Pro Vai Asn Thr Thr Glu Asp Giu Thr Ala Gin IIIe Pro Ala Glu Ala Vai ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG > iie Giy Tyr Leu Asp Leu Giu Giy Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn AGC ACA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT Ser Thr Asn Asn Gly Leu Leu Phe IIe Asn Thr Thr IIe Ala Ser IIe Ala Ala Lys KPI(-4-57; M15A, S17W) Xbal ▶Giu Giu Giy Val Ser Leu Asp Lys Arg Giu Val Val Arg Giu Val Cys Ser Giu Gin RsrII GCT GAG ACC GGT CCG TGC CGT GCA GCT ATC TGG CGC TGG TAC TTT GAC GTC ACT GAA CGA CTC TGG CCA GGC ACG GCA CGT CGA TAG ACC GCG ACC ATG AAA CTG CAG TGA CTT Ala Giu Thr Gly Pro Cys Arg Ala Ala He Trp Arg Trp Tyr Phe Asp Val Thr Glu CGT ANG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC AAC CGT AAC AAC TTT GAC CCA TTC ACG CGA GGT AAG AAA ATG CCG CCA ACG CCG CCG TTG GCA TTG TTG AAA CTG ▶Gly Lys Cys Ala Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp BamHl HindIII ACT GAA GAG TAC TOC ATG GCA GTG TOC GGA TCC GCT ATT TAA GCT T TGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A ▶Thr Glu Glu Tyr Cys Met Ala Val Cys Gly Ser Ala lle

FIG. II

8 1 2 GTECGTGCCGTGCAGCIATCIGGCGCTGGTACTTTGACGT GCACGGCACGTEGATAGACEGCGACCATGAAAC 8 1 3	pTW6165 KPI(-4-57; M15A, S17F)
8 1 4 GTCCGTGCCGTGCAGCTATCTACCGCTGGTACTTTGACGT GCACGGCACG	pTW6166 KPI(-4-57; M15A, S17Y)
8 6 7 GTCCGTGCCGTGCATIGATCIICCGCTGGTACTTTGACGT GCACGGCACGTAACTAGAAGGCGACCATGAAAC 8 6 8	pTW6175 KPI(-4-57; M15L, S17F)
1493 GTCCGTGCCGTGCATIGATCIACCGCTGGTACTTTGACGT GCACGGCACGTAACTAGAIGGCGACCATGAAAC 1494	pBG028 KPI(-4-57; M15L, S17Y)
925 GTCCGTGCCGTGCAATG <u>CACTTC</u> CGCTGGTACTTTGAEGT GCACGGCACGTTAC <u>GTGAAA</u> GGCGACCATGAAAC 926	pTW6183 KPI(-4-57; I16H, S17F)
927 GTCCGTGCCGTGCAATGCACTACCGCTGGTACTTTGACGT GCACCGCACGTTACCTGATGCCGACCATGAAAC 928	pTW6184 KPI(-4-57; I16H, S17Y)
929 GTECGTGCCGTGCAATG <u>CACTGCGCTGCTACTTTGACGT</u> GCACGGCACGTTAC <u>GTGACCGCGACCATGAAAC</u> 930	pTW6185 KPI(-4-57; I16H, S17W)
863 gtccgtgccgtgcagcifactcccgctggtactttgacgt gcacggcacgtfcagtgagggcgaccatgaaac 864	pTW6173 KPI(-4-57; M15A, I16H)
8 6 5 GTCCGTGCCGTGCATIGCACTCCCGCTGGTACTTTGACGT GCACGGCACGTAACCTGAGGGCGACCATGAAAC 8 6 6	pTW6174 KP!(-4-57; M15L, I16H)

pTW 6166

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α-factor
  ATG AGA TIT CCT TCA ATT TIT ACT GCA GIT TIA TIC GCA GCA TCC TCC GCA TIA GCT
  THE TET ANA GGA AGT TAN ANN TGN CGT CAN ANT ANG CGT CGT AGG AGG CGT ANT CGN
▶Met Arg Phe Pro Ser lie Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala
  GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG
 Ala Pro Val Asn Thr Thr Glu Asp Glu Thr Ala Gin ile Pro Ala Glu Ala Val
  ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG
File Gly Tyr Leu Asp Leu Glu Gly Asp Phe Asp Val Aig Val Leu Pro Phe Ser Asn
 AGC ACA MAT MAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT
Ser Thr Asn Asn Gly Leu Leu Phe lie Asn Thr Thr lie Ala Ser lie Ala Ala Lays
                                                     KP!(-4-57; M15A, S17Y)
 GAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA CTT CTT CCC CAT AGA GAT CTA TTT TCT CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT
FGIU GIU GIY Val Ser Leu Asp Lys Arg GIU Val Val Arg GIU Val Cys Ser GIU GIN
                Rsrll
             Agel
 CCT GAG ACC GGT CCG TGC CGT CCA GCT ATC TAC CGC TGG TAC TTT GAC GTC ACT GAA CGA CTC TGG CCA GGC ACG CGT CGA TAG ATG CGA ACC ATG AAA CTG CAG TGA CTT
Ala Giu Thr Gly Pro Cys Arg Ala Ala ile Tyr Arg Trp Tyr Phe Asp Val Thr Giu
 GGT ANG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC ANC CGT ANC ANC TTT GAC CCA TTC ACG CGA GGT ANG ANA ATG CCG CCA ACG CCG CTG GCA TTG GTA ANA CTG
FGIV Lys Cys Ala Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp
                                                   BamH!
 ACT GAA GAG TAC TGC ATG GCA GTG TGC GGA TCC GCT ATT TAA GCT TTGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A
Thr Glu Glu Tyr Cys Met Ala Val Cys Gly Ser Ala ile
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α-factor
 ATG AGA TIT CCT TCA ATT TIT ACT GCA GIT TTA TTC GCA GCA TCC TCC GCA TTA GCT TAC TCT AAA GGA AGT TAA AAA TGA CGT CAA AAT AAG CGT CGT AGG AGG CGT AAT CGA
Met Arg Phe Pro Ser ile Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala
 GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG
PAla Pro Val Asn Thr Thr Thr Giu Asp Giu Thr Ala Gin He Pro Ala Giu Ala Val
 ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC
  TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG
Flie Gly Tyr Leu Asp Leu Glu Gly Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn
AGC ACA AAT AAC GGG TTA TTG TIT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TCT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT Ser Thr Asn Asn Gly Leu Leu Phe lie Asn Thr Thr lie Ala Ser lie Ala Ala Lys
                                                  KPI(-4-57; M15L, S17F)
                         Xbal
 GAA GAA GOG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA
 CTT CTT CCC CAT AGA GAT CTA TIT TCT CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT
▶Giu Giu Giy Val Ser Leu Asp Lys Arg Giu Val Val Arg Giu Val Cys Ser Giu Gin
               Rsrll
                                                                                 Aatll
            Agel
 GCT GAG ACC GGT CCG TGC CGT GCA TTG ATC TTC CGC TGG TAC TTT. GAC GTC ACT GAA CGA CTC TGG CCA GGC ACG GCA CGT AAC TAG AAG GCG ACC ATG AAA CTG CAG TGA CTT
Ala Giu Thr Gly Pro Cys Arg Ala Leu IIe Phe Arg Trp Tyr Phe Asp Val Thr Glu
 GGT ANG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC ANC CGT ANC ANC TTT GAC CCA TTC ACG CGA GGT ANG ANA ATG CCG CCA ACG CCG CCG TTG GCA TTG TTG ANA CTG
FGIY Lys Cys Ala Pro Phe Phe Tyr Giy Giy Cys Giy Giy Asn Arg Asn Asn Phe Asp
                                                  BamHI
                                                                       Hindll
 ACT GAA GAG TAC TOO ATG OCA GTG TOO OGA TOO GCT ATT TAA GCT T
 TGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A
▶Thr Giu Giu Tyr Cys Met Ala Val Cys Gly Ser Ala lie
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α-factor
 ATG AGA TIT CCT TCA ATT TIT ACT GCA GIT TIA TIC GCA GCA TCC TCC GCA TIA GCT
 TAC TOT AAA GGA AGT TAA AAA TGA CGT CAA AAT AAG CGT CGT AGG AGG CGT AAT CGA
Met Arg Phe Pro Ser IIe Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala
 GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC
 CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG
▶Ala Pro Val Asn Thr Thr Glu Asp Glu Thr Ala Gln IIe Pro Ala Glu Ala Val
 ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG
▶ He Gly Tyr Leu Asp Leu Glu Gly Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn
 AGC ACA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT
Ser Thr Asn Asn Gly Leu Leu Phe lie Asn Thr Thr lie Ala Ser lie Ala Ala Lys
                                                 KPI(-4-57; M15L, S17Y)
                         Xbal
 GAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA CTT CTT CCC CAT AGA GAT CTA TTT TCT CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT
FGIU GIU GIY VAI Ser Leu Asp Lys Arg Giu Vai Vai Arg Giu Vai Cys Ser Giu Gin
              Rsril
            Agel
                                                                                Aatil
 GCT GAG ACC GGT CCG TGC CGT GCA TTG ATC TAC CGC TGG TAC TTT GAC GTC ACT GAA CGA CTC TGG CCA GGC ACG GCA CGT AAC TAG ATG GCG ACC ATG AAA CTG CAG TGA CTT
Ala Glu Thr Gly Pro Cys Arg Ala Leu IIe Tyr Arg Trp Tyr Phe Asp Val Thr Glu
 GGT AAG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC AAC CGT AAC AAC TTT GAC CCA TTC ACG CGA GGT AAG AAA ATG CCG CCA ACG CCG CTG GCA TTG TTG AAA CTG
Fig. Lys Cys Ala Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp
                                                 BamHI
                                                                      HindIII
 ACT GAA GAG TAC TOC ATG GCA GTG TOC GGA TCC GCT ATT TAA GCT TTGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A
Thr Giu Giu Tyr Cys Met Ala Val Cys Giy Ser Ala ile
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FIG. 15

α-factor ATG AGA TIT CCT TCA ATT TIT ACT GCA GTT TTA TTC GCA GCA TCC TCC GCA TTA GCT TAC TCT AAA GGA AGT TAA AAA TGA CGT CAA AAT AAG CGT CGT AGG AGG CGT AAT CGA ▶Met Arg Phe Pro Ser IIe Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG PAIR Pro Val Asn Thr Thr Thr Glu Asp Giu Thr Ala Gin II. Pro Ala Glu Ala Val ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG ≯ile Giy Tyr Leu Asp Leu Giu Giy Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn AGC ACA AAT AAC GOG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT Ser Thr Asn Asn Gly Lou Lou Pho II o Asn Thr Thr II o Ala Ser II o Ala Ala Lys KPI(-4-57; I16H, S17F) Xbal GAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA CTT CTT CCC CAT AGA GAT CTA TTT TCT CTC CAA CAA TCT CTC CAC AGA CTT GTT ▶Giu Giu Giy Vai Ser Leu Asp Lys Arg Giu Vai Vai Arg Giu Vai Cys Ser Giu Gin Rsrll Agel CCT GAG ACC GGT CCG TGC CGT GCA ATG CAC TTC CGC TGG TAC TTT GAC GTC ACT GAA CGA CTC TGG CCA GGC ACG GCA CGT TAC GTG AAG GGG ACC ATG AAA CTG CAG TGA CTT >Ala Glu Thr Gly Pro Cys Arg Ala Met His Phe Arg Trp Tyr Phe Asp Val Thr Glu GGT AAG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC AAC CGT AAC AAC TTT GAC CCA TTC ACG CGA GGT AAG AAA ATG CCG CCA ACG CCG TTG GCA TTG TTG AAA CTG FGIY Lys Cys Ala Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp BamHI HindIII ACT GAA GAG TAC TOC ATG GCA GTG TGC GGA TCC GCT ATT TAA GCT T TGA CIT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A ▶Thr Giu Glu Tyr Cya Met Ala Val Cys Gly Ser Ala ile

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α-factor
  ATG AGA TIT CCT TCA ATT TIT ACT GCA GTT TTA TTC GCA GCA TCC TCC GCA TTA GCT
 TAC TOT ANA GGA AGT TAN ANA TGA CGT CAN ANT ANG CGT CGT AGG AGG CGT ANT CGA
▶Met Arg Phe Pro Ser ile Phe Thr Aia Vai Leu Phe Ala Aia Ser Ser Ala Leu Ala
 GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG
PALA Pro Val Asn Thr Thr Thr Glu Asp Glu Thr Ala Gin Ile Pro Ala Glu Ala Val
 ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC
 TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG
▶ Ite Giy Tyr Leu Asp Leu Giu Giy Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn
 AGC ACA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT
Ser Thr Asn Asn Gly Leu Leu Phe ile Asn Thr Thr ile Ala Ser lie Ala Ala Lys
                                                 KPI(-4-57; 116H, S17Y)
                         Xbal
 GAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA CTT CTT CCC CAT AGA GAT CTA TTT TCT CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT
▶Giu Giu Giy Val Ser Leu Asp Lys Arg Giu Val Val Arg Giu Val Cys Ser Giu Gin
               Rsrll
            Agel
                                                                               Aatll
 GCT GAG ACC GGT CCG TGC CGT GCA ATG CAC TAC CGC TGG TAC TTT GAC GTC ACT GAA CGA CTC TGG CCA GGC ACG GCA CGT TAC GTG ATG GCG ACC ATG ANA CTG CAG TGA CTT
PAIR GIU Thr Gly Pro Cys Arg Ala Met His Tyr Arg Trp Tyr Phe Asp Val Thr Glu
 GGT AAG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC AAC CGT AAC AAC TTT GAC CCA TTC ACG CGA GGT AAG AAA ATG CCG CCA ACG CCG CCG TTG GCA TTG TTG AAA CTG
Bigly Lys Cys Ala Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp
                                                 BamHi
 ACT GAA GAG TAC TGC ATG GCA GTG TGC GGA TCC GCT ATT TAA GCT T TGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A
▶Thr Glu Glu Tyr Cys Met Ala Val Cys Gly Ser Ala ile
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FIG. 17

α-factor ATG AGA TIT CCT TCA ATT TIT ACT GCA CTT TTA TTC GCA GCA TCC TCC GCA TTA GCT TAC TOT ANA GGA AGT TAN ANA TGA CGT CAN ANT ANG CGT CGT AGG AGG CGT ANT CGA PMet Arg Phe Pro Ser ite Phe Thr Ala Val Leu Phe Ata Ata Ser Ser Ala Leu Ata GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG PALE Pro Val Asn Thr Thr Thr Glu Asp Glu Thr Ala Gin Ile Pro Ala Glu Ala Val ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG File Gly Tyr Leu Asp Leu Glu Gly Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn AGC ACA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT Ser Thr Asn Asn Gly Leu Leu Phe II.e Asn Thr Thr II.e Ala Ser II.e Ala Ala Lys KPI(-4-57; I16H, S17W) GAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA CTT CTT CCC CAT AGA GAT CTA TTT TCT CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT GTU GIU GIU GIU GIU Vai Vai Vai Vai Vai Cys Ser Giu Gin Xbal Rsrll Agel GCT GAG ACC GGT CCG TGC CGT GCA ATG CAC TGG CGC TGG TAC TTT GAC GTC ACT GAA CGG CTC TGG CCA GGC ACG GCA CGT TAC GTG ACC GCG ACC ATG AAA CTG CAG TGA CTT PAIA GIU Thr Gly Pro Cys Arg Ala Met His Trp Arg Trp Tyr Phe Asp Val Thr Giu GOT ANG TOC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC AAC CGT AAC AAC TTT GAC CCA TTC ACG CGA GGT AAG AAA ATG CCG CCA ACG CCG CCG TTG GCA TTG TTG AAA CTG FGIY Lys Cys Ala Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp BamHI ACT GAA GAG TAC TOC ATG GCA GTG TOC GGA TCC GCT ATT TAA GCT T TGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A ▶Thr Glu Glu Tyr Cys Met Ala Val Cys Gly Ser Ala ile

FIG. 18

α-factor ATG AGA TIT CCT TOA ATT TIT ACT GOA GIT TIA TIC GOA GOA TOO TOO GOA TIA GCT TAC TCT AAA GGA AGT TAA AAA TGA CGT CAA AAT AAG CGT CGT AGG AGG CGT AAT CGA ▶Met Arg Phe Pro Ser ite Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG PAla Pro Vai Asn Thr Thr Thr Giu Asp Giu Thr Ala Gin IIe Pro Ala Giu Ala Val ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG File Gly Tyr Leu Asp Leu Glu Gly Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn AGC ACA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT ▶ Ser Thr Asn Asn Gly Leu Leu Phe IIe Asn Thr Thr IIe Ala Ser IIe Ala Ala Lys KPI(-4-57; M15A, I16H) Xbal GAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA
CTT CTT CCC CAT AGA GAT CTA TTT TCT
CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT ⊳Giu Giu Giy Val Ser Leu Asp Lys Arg Giu Val Val Arg Giu Val Cys Ser Giu Gin RsrII Aatii Agel CCT CAG ACC CGT CCG TGC CGT GCA CCT CAC TCC CGC TGG TAC TTT GAC GTC ACT GAA CCG. CTC TGG CCA GGC ACG GCA CGT CGA GTG AGG GCA ACG ACG ACG ACG ACG ACG AAA CTG CAG TGA CTT Ala Glu Thr Gly Pro Cys Arg Ala Ala His Ser Arg Trp Tyr Phe Asp Val Thr Glu GOT ANG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC AAC CGT AAC AAC TTT GAC CCA TTC ACG CGA GGT ANG ANN ATG CCG CCA ACG CCG CTG GCA TTG TTG ANN CTG FGIY Lys Cys Ala Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp BamHI ACT GAA GAG TAC TOC ATG GCA GTG TGC GGA TCC GCT ATT TAA GCT T TGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A ▶Thr Giu Giu Tyr Cys Met Ala Val Cys Gly Ser Ala ile

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α-factor
 ATG AGA TIT CCT TCA ATT TIT ACT GCA GTT TTA TTC GCA GCA TCC TCC GCA TTA GCT TAC TCT AAA GGA AGT TAA AAA TGA CGT CAA AAT AAG CGT CGT ACG ACG CGT AAT CGA
Met Arg Phe Pro Ser ile Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala
 GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG. TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG
PAIR Pro Val Asn Thr Thr Glu Asp Glu Thr Ala Gin He Pro Ala Glu Ala Val
 ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG
▶ He Gly Tyr Leu Asp Leu Glu Gly Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn
 AGC ACA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT
▶ Ser Thr Asn Asn Gly Leu Leu Phe ile Asn Thr Thr lie Ala Ser lie Ala Ala Lys
                                                        KPI(-4-57; M15L, I16H)
                            Xbal
 CAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA CTT CTT CTC CAC AGA GAT CTA TTT TCT CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT
▶Glu Glu Gly Val Ser Leu Asp Lys Arg Glu Val Val Arg Glu Val Cys Ser Glu Gin
                 RsrII
 GCT GAG ACC GGT CCG TGC CGT GCA TTG CAC TCC CGC TGG TAC TTT GAC GTC ACT GAA CGG CTC TGG CCA GGC ACG GCA CGT AAC GTC AGG GCA CTT AAA CTG CAG TGA CTT
              Agel
PAIA GIU Thr Gly Pro Cys. Arg Ala Leu His Ser Arg Trp Tyr Phe Asp Val Thr Glu
  GGT ANG TOC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC ANC CGT ANC ANC TTT GAC CCA TTC ACG CGA GGT ANG ANA ATG CCG CCA ACG CCG CTG TTG GCA TTG TTG ANA CTG
FGIY LYS Cys Als Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp
                                                                               HindIII
                                                        BamHI
  ACT GAA GAG TAC TOC ATG OCA GTG TOC GGA TCC OCT ATT TAA OCT T
TGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A
▶Thr Glu Glu Tyr Cys Met Ala Val Cys Gly Ser Ala IIe
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FIG. 20

KPI(-4-57; M15A, S17W) TW6165

FIG. 21

KPI(-4-57; M15A, S17Y) TW6166

FIG. 22

KPI(-4-57; M15L, S17F) TW6175

FIG. 23

KPI(-4-57; M15L, S17Y) BG028

FIG. 24

KPI(-4-57; I16H, S17F) TW6183

FIG. 25

KPI(-4-57; I16H, S17Y) TW6184

FIG. 26

KPI(-4-57; I16H, S17W) TW6185

FIG. 27

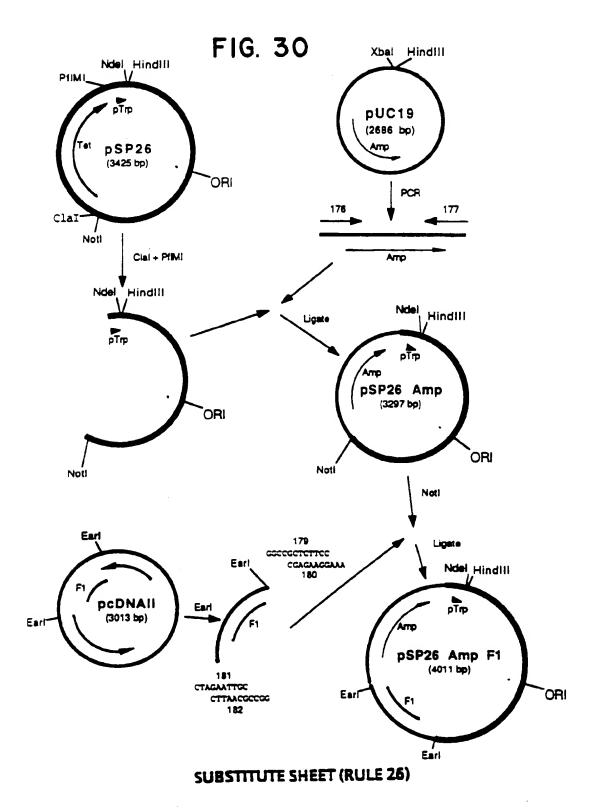
KPI(-4-57; M15A, S17F) DD185

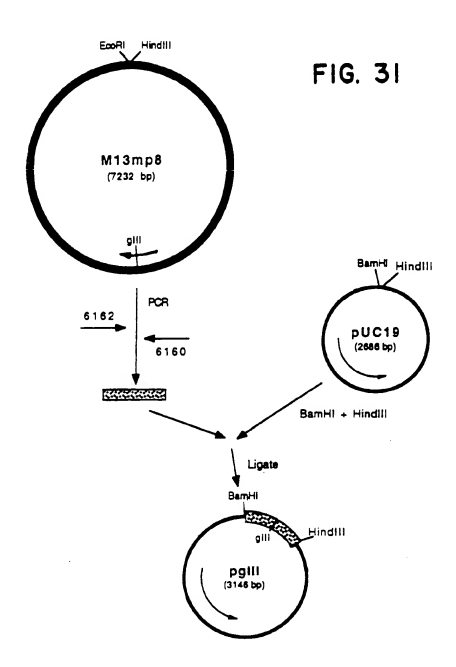
FIG. 28

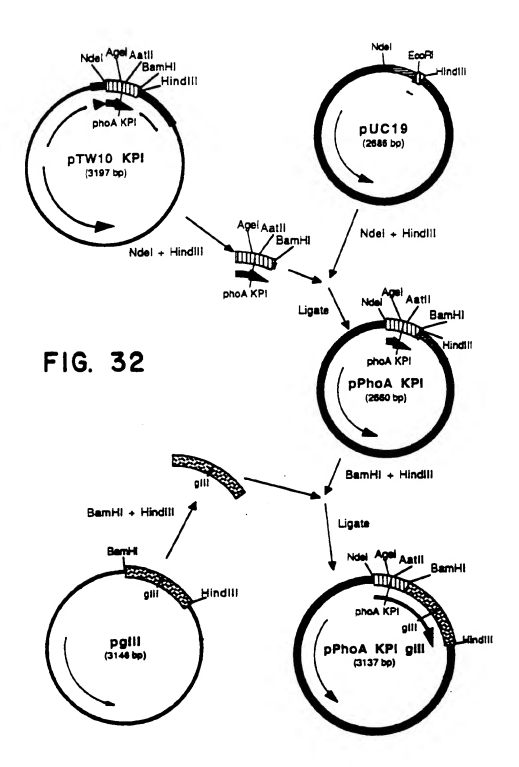
KPI(-4-57; M15A, I16H) TW6173

FIG. 29

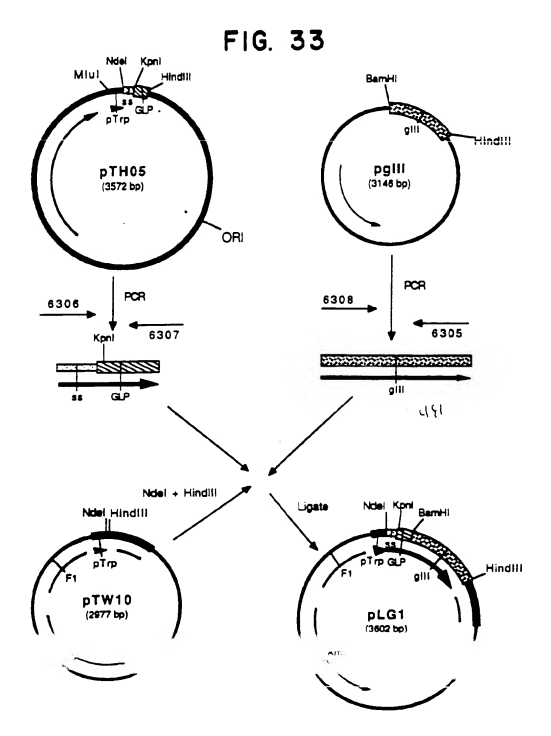
KPI(-4-57; M15L, I16H) TW6174





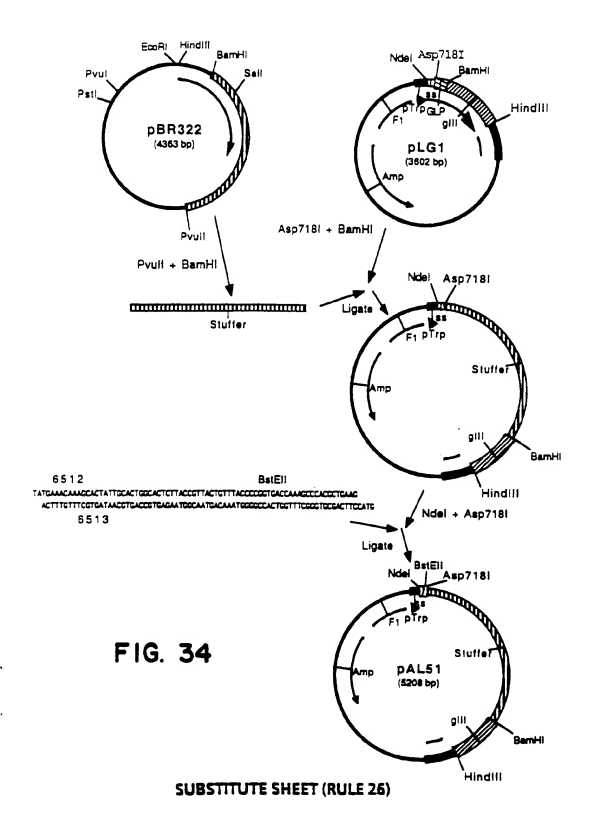


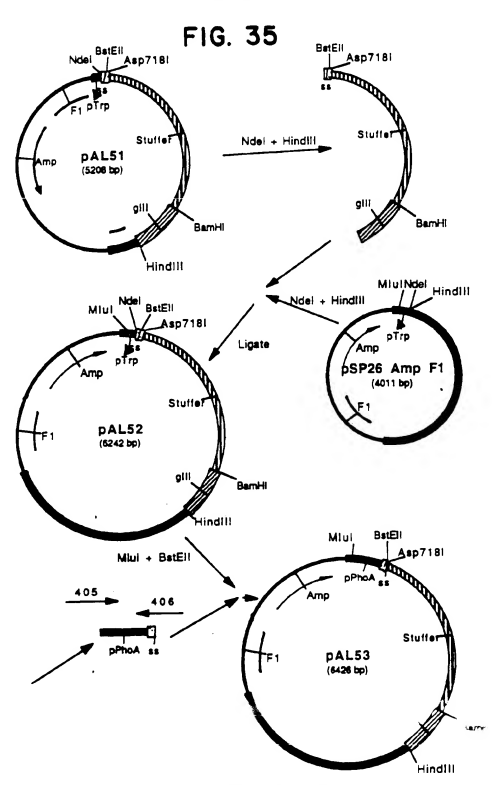
SUBSTITUTE SHEET (RULE 26)



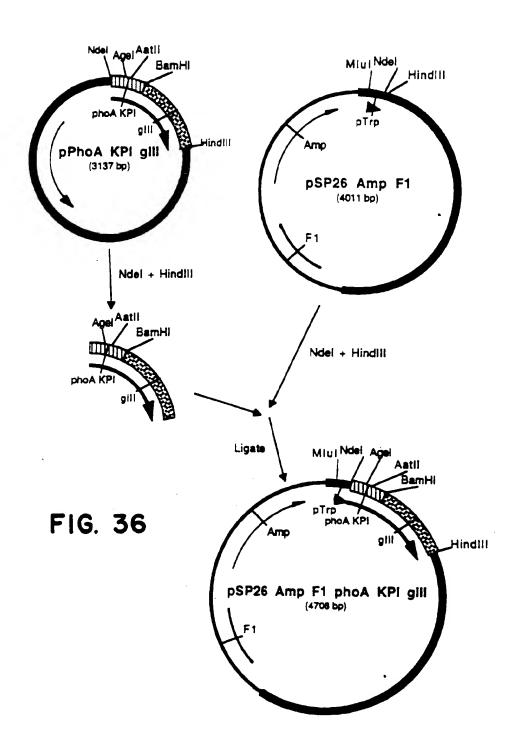
SUBSTITUTE SHEET (RULE 26)

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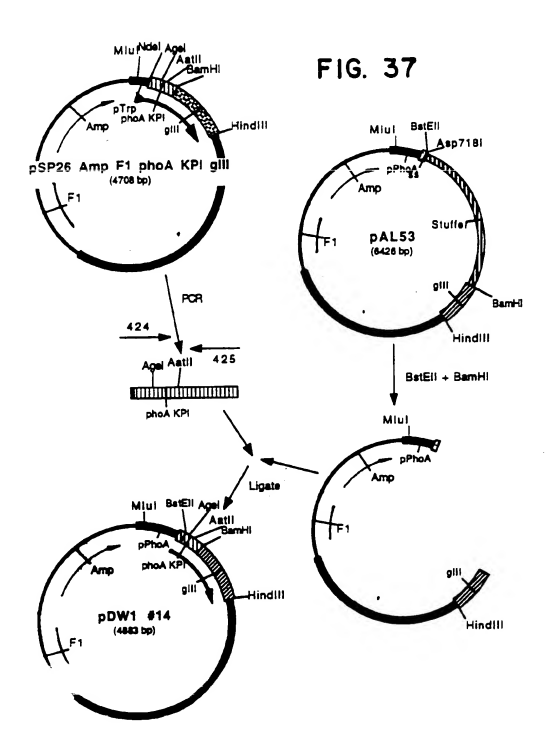




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SUBSTITUTE SHEET (RULE 26)

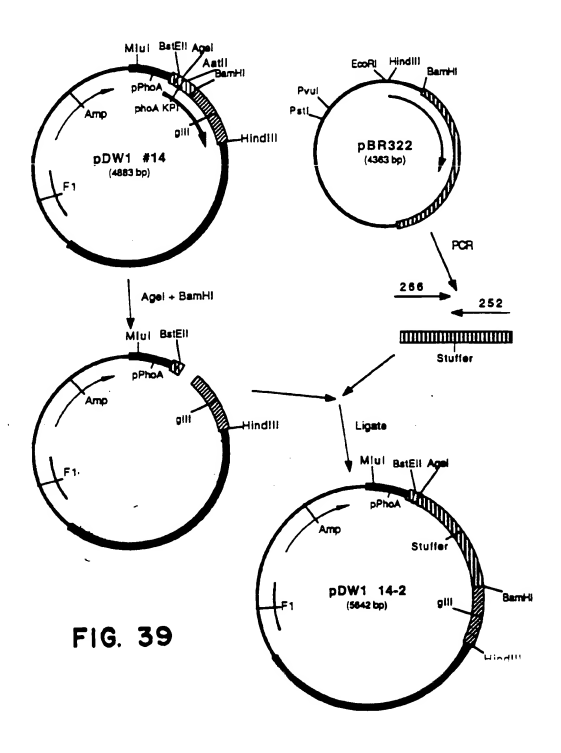
FIG. 38

phoA signal GTG AAA CAA AGC ACT ATT GCA CTG GCA CTC TTA CCG TTA CTG TTT ACC CCG GTG ACC AAA ▶ Val Lys Gin Ser Thr lie Ala Leu Ala Leu Leu Pro Leu Leu Phe Thr Pro Val Thr Lys KPI (1-55)

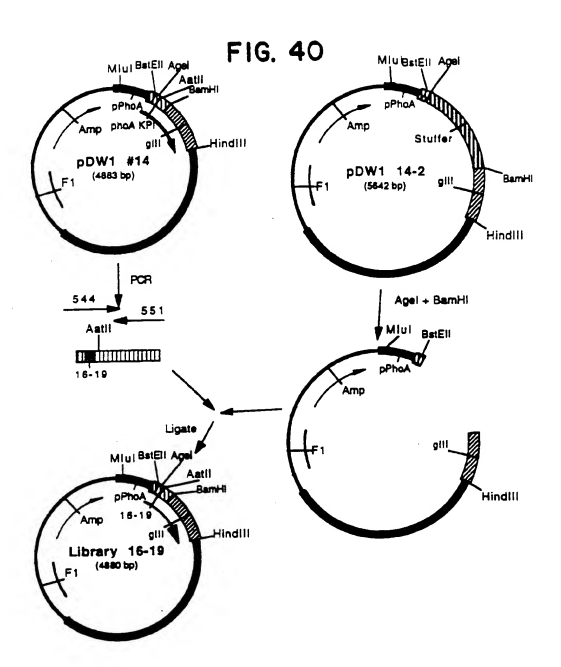
Agel

GCC GAG GTG TGC TCT GAA CAA GCT GAG ACC GGT CCG TGC CGT GCA ATC ATC TCC CGC TGC PAIA Glu Val Cys Ser Glu Gin Ala Glu Thr Gly Pro Cys Arg Ala Met Ile Ser Arg Tro Aatii TAC TIT GAC GTC ACT GAA GGT AAG TGC GCT CCA TTC TIT TAC GGC GGT TGC GGC GAC Tyr Phe Asp Val Thr Glu Gly Lys Cys Ala Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn BamH1 CGT AAC AAC TIT GAC ACT GAA GAG TAC TGC ATG GCA GTG TGC GGA TCC GGT GGT, GGC TCT Arg Asn Asn Phe Asp Thr Glu Glu Tyr Cys Met Ala Val Cys Gly Ser Gly Gly Gly Ser GGT TCC GGT GAT TIT GAT TAT GAA AAG ATG GCA AAC GCT AAT AAG GGG GCT ATG ACC GAA ▶Gly Ser Gly Asp Phe Asp Tyr Glu Lys Met Ala Asn Ala Asn Lys Gly Ala Met Thr Glu AAT GCC GAT GAA AAC GCG CTA CAG TCT GAC GCT AAA GGC AAA CTT GAT TCT GTC GCT ACT ≯Asn Ala Asp Giu Asn Ala Leu Gin Ser Asp Ala Lys Giy Lys Leu Asp Ser Val Ala Thr GAT TAC GGT GCT GCT ATC GAT GGT TTC ATT GGT GAC GTT TCC GGC CTT GCT AAT GGT AAT ≯Asp Tyr Giy Ala Ala ile Asp Giy Phe ile Giy Asp Val Ser Giy Leu Ala Ash Giy Ash GGT GCT ACT GGT GAT TIT GCT GGC TCT AAT TCC CAA ATG GCT CAA GTC GGT GAC GGT GAT FGIY Ala Thr Gly Asp Phe Ala Gly Ser Asn Ser Gln Met Ala Gln Val Gly Asp Gly Asp AAT TOA COT TTA ATG AAT AAT TTC CGT CAA TAT TTA CCT TCC CTC CCT CAA TCG GTT GAA PASN Ser Pro Leu Met Asn Asn Phe Arg Gin Tyr Leu Pro Ser Leu Pro Gin Ser Val Glu TOT COC CCT TIT GIC TIT GOC GCT GGT ANA CCA TAC GAA TIT TCT ATT GAT TGT GAC ANA > Cys Arg Pro Phe Val Phe Gly Ala Gly Lys Pro Tyr Glu Phe Ser Ile Asp Cys Asp Lys ATA AAC TTA TTC CGT GGT GTC TTT GCG TTT CTT TTA TAT GTT GCC ACC TTT ATG TAT GTA File Asn Leu Phe Arg Gly Val Phe Ala Phe Leu Leu Tyr Val Ala Thr Phe Met Tyr Val TIT TOT ACE TIT COT AAC ATA CTG CGT AAT AAG GAG TOT TAA TA Phe Ser The Phe Ala Asn ile Leu Arg Asn Lys Giu Ser ...

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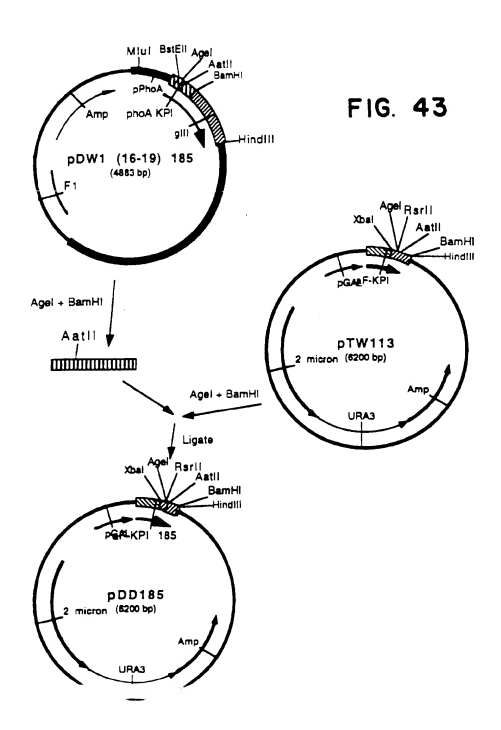
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phoA signal				BstEll
CTC ANA CAN AGC	ACT ATT GCA	CTG GCA CTC	TTA CCG TTA CT	G TTT ACC CCG GTG ACC AAA
Val Lys Gin Ser	Thr lie Ala	Leu Ala Leu	Leu Pro Leu Le	u Phe Thr Pro Val Thr Lys
KPI (1-55;	16-19)	Δαρ		16-19
		~~ ~~ ~~	רכיד ררכי זיכר רכי	T NNS NNS NNS NNS TGG TAC
FAIR Glu Val Cys	Ser Glu Gin	Ala Glu Thr	Gly Pro Cys A	g ??? ??? ??? ??? Trp Tyr
•				
Aatii	C12 CCT 11C	TO COT CCA	TTC TTT TAC GO	SC GGT TGC GGC GGC AAC CGT
bone Asp Val The	GIU GIV LVS	Cvs Ala Pro	Pho Pho lyr Gi	y Giy Cys Giy Giy Asii Aiy
THE ASP VET III.	2.0 2.7 2,	•	0.	gill A TCC GGT GGT GGC TCT GGT A SA GUY GUY GUY SAF GUY
	.cm (3) (3)	TAC TOC ATG	GCA GTG TGC GG	A TCC GGT GGT GGC TCT GGT
AAC AAC TTT GAC	Thr Glu Glu	Tyr Cys Met	Ala Val Cys Gi	y Ser Gly Gly Gly Ser Gly
TCC GGT GAT TTT	GAT TAT GAA	AAG ATG GLA	Asn Ala Asn L	AG GGG GCT ATG ACC GAA AAT /s Gly Ala Met Thr Glu Asn
GCC GAT GAA AAC	GCG CTA CAG	TCT GAC GCT	ANN GOC ANN C	TT GAT TOT GTC GCT ACT GAT
				au Asp Ser Val Ala Thr Asp
TAC GGT GCT GCT	ATC GAT GGT	TTC ATT GGT	GAC GITT TCC G	SC CTT GCT AAT GGT AAT GGT
FTyr Gly Ala Ala	ile Asp Gly	Phe lie Gly	Asp Val Ser G	ly Leu Ala Asn Gly Asn Gly
			gi	11
GCT ACT GGT GAT	TTT GCT GGC	TCT AAT TCC	CAA ATG OCT C	AA GTC GGT GAC GGT GAT AAT
►Ala Thr Gly Asp	Phe Ala Gly	Ser Asn Ser	Gin Met Ala G	In Val Gly Asp Gly Asp Asn
TO COT THE ATG	AAT AAT TTC	CGT CAA TAT	TTA CCT TCC C	TO COT CAN TOG GTT GAN TGT
≯Ser Pro Leu Met	Asn Asn Phe	Arg Gin Ty	r Leu Pro Ser L	eu Pro Gin Ser Val Glu Cys
	- سسة فحد فحية	COT ANA CC	TAC GAA TIT T	CT ATT GAT TGT GAC AAA ATA
►Ara Pro Phe Val	Phe Gly Ala	Gly Lys Pro	Tyr Glu Phe S	er lie Asp Cys Asp Lys ile
-				
AAC TTA TTC CGT	COT GIC TIL	Ala Pha Lau	Leu Tyr Val A	CC ACC TTT ATG TAT GTA TTT is Thr Phs Met Tyr Val Phe
TOT ACG TIT GOT	AAC ATA CTO	CGT AAT AA	GAG TCT TAA T	'A
Ser Thr Phe Als	AST ITE LEV	Arg Asn Ly	S Call 341	

ph	oA s	igna	ll .		_											8	tEII		
GTG	AAA	CAA	AGC	ACT	ATT	GCA	CTG	GCA	CTC	TTA	೦೦೦	TTA	CTG	TTT	ACC	೦೦೦	GTG	ACC	AAA
P Val	Lys	GIn	Ser	Thr	11•	Ala	Leu	Ala	Leu	Leu	Pro	Leu	Leu	Phe	Thr	Pro	Va I	Thr	Lys
				M15			F۱												
ccc								GAG		CCT	ccc	TGC	CCT	GCA	GCT.	ATC	TTC	CGC	TGG
GCC: ►Ala	GAG	Vel	Cve	Ser	GLU	Glo	Ala	Glu	Thr	GIV	Pro	Cvs	Ara	Al a	Ala	110	Phe	Arg	Trp
PAIR	Giu	V & 1	U ,.	.	<u> </u>	—		•		- ,		-,-	•						•
		Aatl	١																
TAC	TIT	CAC	GTC	ACT	GAA	GGT	AAG	TGC	GCT	CCY	TTC	TIT	TAC	GGC	GGT	ICC	œc	<u>@</u>	AAC
≯ Tyr	Ph e	Asp	Va I	Thr	GI U	GI y	Lys	Cys	Ala	Pro	Ph e	Phe	Tyr	Gal y				GI y	ASN
														Bami	- 11	g	<u> </u>		
CGT	AAC	AAC	TIT	GAC	ACT	GAA	GAG	TAC	TGC	ATG	GCA	GTG	TGC	GGA	TCC	GGT	GGT	œ	TCT
PArg	Asn	Asn	Phe	Asp	Thr	GI u	GI u	Tyr	Cys	Me !	Ala	Val	Cys	GI y	Ser	GI y	GI y	GI y	Ser
GGT		~~	CAM	-	CAT	ጥልጥ	GAA	AAG	ATC	GCA.	AAC	CCT	AAT	AAG	GGG	CCT	ATG	ACC	GAA
GGT ▶ GI y	TCC	GUT	Aan	Phe	Aso	Tvr	Glu	Lys	Met	Ala	Asn	Ala	Asn	Lys	Gly	Ala	Met	Thr	Glu
-																			
AAT	CCC	GAT	CYY	AAC	CCC	CTA	CAG	TCT	GAC	CCT	AAA	œc	AAA	CTT	CAT	TCT	GTC	GCT	ACT
≯Asn	Ala	Asp	Gi u	Asn	Ala	Leu	Gin	Ser	ASP	AIR	Lys	Car y	Lys	Leu	ABP	307	VE:	A1 =	1131
GAT	TAC	GGT	GCT	GCT	ATC	GAT	GGT	TTC	ATT	GGT	GAC	GIT	TCC	GGC	CIT	CCI	AAT	GGT	AAT
≯Asp	Tyr	GI y	AIR	Al s	110	Asp	GI y	Phe	1 i •	Gi y	Asp	Val	Ser	GI y	Leu	A la	Asn	GI y	Asn
GGT			~~	C) E	-	CCT1	ccc		AAT	TCC	CAA	ATC	ركيات	CAA	CTC.	CCT	GAC	CCT	GAT
GGT ▶ GIy	GCT	ACT	GUT	Asn	Phe	Ala	Giv	Ser	Asn	Ser	GIA	Me 1	Ala	Gin	Val	Gly	Asp	Gly	Asp
-																			
AAT	TCA	CCI	TTA	ATG	AAT	AAT	TTC	CCL	CAA	TAT	TTA	CCI	TCC	crc	CCT	CAA	TCG	GIT	GAA
►Asn	Ser	Pro	Leu	Me	Asn	ASD	PRE	Arg	GE N	ıyr	Leu	Pro	30 f	Leu	FIQ	Gin	301	V	Gi U
TGT	CGC	CCT	TTT	GTC	TIT	GGC	GCT	GCT	AAA	CCA	TAC	GAA	TTT	TCT	ATT	GAT	TGT	GAC	AAA
→ Cys	A rg	Pro	Phe	Val	Phe	GI y	Al a	GI y	Lys	Pro	Tyr	GI u	Phe	Ser	110	Asp	Cys	Asp	Lys
ATA					~~	~~~	-	~~	Teres	مبعثت	TTTA	ጥልጥ	بتعلث	~~	300	Teleb	ATC:	ጥልጥ	CTA
ATA 110	AAC	TTA	TTC	Arn	GUT	Val	Phe	Ala	Phe	Leu	Leu	TVI	Val	Ala	Thr	Phe	Met	Tyr	Val
															,		- •	,	
TTT	TCT	ACG	TTT	GCT	AAC	ATA	CIG	CCI	AAT	AAG	CAG	TCT	TAA	TA					
▶ Ph e	Ser	Thr	Phe	Al a	Asn	110	Leu	Arg	Asn	Lys	GIU	Ser							



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α-factor
 ATG AGA TIT CCT TCA ATT TIT ACT GCA GIT TIA TIC GCA GCA TCC TCC GCA TTA GCT
 TAC TOT AAA GGA AGT TAA AAA TGA CGT CAA AAT AAG CGT CGT AGG AGG CGT AAT CGA
>Met Arg Phe Pro Ser lie Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala
GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG
PALE Pro Val Asn Thr Thr Thr Glu Asp Glu Thr Ala Gln Ile Pro Ala Glu Ala Val
 ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG
▶ II e Giy Tyr Leu Asp Leu Giu Giy Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn
AGC ACA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT
≯Ser Thr Asn Asn Gly Leu Leu Phe lie Asn Thr Thr lie Ala Ser lie Ala Ala Lys
                                                  KP1(-4-57; M15A, S17F)
                         Xbal
 GAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA CTT CTT CCC CAT AGA GAT CTA TTT TCT CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT
▶Glu Glu Gly Val Ser Leu Asp Lys Arg Glu Val Val Arg Glu Val Cys Ser Glu Gin
               RsrII
             Agel
 GCT GAG ACC GGT CCG TGC CGT GCA CCT ATC TTC CGC TGG TAC TTT GAC GTC ACT GAA CGA CTC TGG CCA GGC ACG GCA CGT CGA TAG AAG GCG ACC ATG AAA CTG CAG TGA CTT
PAIS GIU Thr Gly Pro Cys Arg Ala Ala II Phe Arg Trp Tyr Phe Asp Val Thr Glu
  GGT ANG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC ANC CGT ANC ANC TTT GAC CCA TTC ACG CGA GGT ANG ANA ATG CCG CCA ACG CCG CCG TTG GCA TTG TTG ANA CTG
FGIY Lys Cys Ala Pro Phe Phe Tyr Giy Giy Cys Giy Giy Asn Arg Asn Asn Phe Asp
                                                  BamH!
 ACT GAA GAG TAC TGC ATG GCA GTG TGC GGA TCC GCT ATT TAA GCT T
 TGA CTT CTC ATG ACG TAC CCT CAC ACG CCT AGG CGA TAA ATT CGA A
 ▶Thr Glu Giu Tyr Cys Met Ala Val Cys Gly Ser Ala lie
```

161.0

3 KPI (-4-57) Variant

196.0 nd nd 56.0 nd 159.0 214.0 473.0 nd nd nd nd nd nd nd od 0.6 0.6

Protease inhibition by KPI (-4-57) variants

			Хį	Kis (nM)					
Yariant	inai	d	3	Substitution	8,		kallikrein	쩟	
ᇫ	KPI (-4-57)	'n	<u>0</u>	<u> </u>	9	ò	45.00	3718.0	
	KBI (4.57: M15A. S17F)		<	ı			0.39	150.0	
	KPI (4-57-M15A S17W)		<	3			0.65	206.0	
	KPI (4.57: M15A, S17Y)		⋖	>			0.40	73.0	
. ע	KPI (4:57: M15)		ر	L			0.50	35.0	
	KPI (4.57-M15) S17Y)		_	>			1.10	93.8	
~	KPI (4-57-116H S17F)			ш Ш			1.20	12440.0	
	KPI (4.57-116H S17V)			≻			0.91	14000.0	
	KDI (4.57-116H S17W)			≯			1.30	388.0	
, ,	KPI (4-57; M15A 116H)		∢	ī			1.00	1432.0	
	(A)		ب	r		•	0.90	2796.0	
r	KPI (4-57: M151 S17Y R18H)		_1	>	ĭ		9.00	19.4	
	KPI (4.57: M15A S17Y B18H)		∢	>	I		0.64	14.5	
	KDI (4.57: T9V M151 S17Y R18H)	>	_	>	I		3.20	6.7	
	KDI (4.57: T9V M15A S17Y, R18H)	>	<	>	I		0.75	5.8	
	KDI (4.57: M45) 116F S17K)		_	T.			7.90	1385.0	
	KDI (4.57: M151 116F S17K G37Y)		_	A X		>	1.10	15640.0	
	KPI (4-57; M15L, 116F, S17K, G37L)		_	Ŧ		_	1.30	7473.0	

15. (165. (1

	FIG 46(1)		Inhibition Ki (nM)	Ki (nM)	
4	Continuo	P. kaili	Plasmin	₫	Я
Vanan	Coquestion Community and Community Research	20.00	0.23	0.0009	
	RPDFCLEPPYTGFCAALLINIFICATION COMPLYCOCRAKSNIFKSAEDCHRTCGGA	0.91	0.17	3983.0	
Aprotinin R15, S42	DPCLEPPTTGPCRAKL LATE INVIDENCE OF THE DEPTCH OF THE THE VICENCE OF THE VICENCE O	45.00	34.00	3718.0	161.0
KPI (-4-57)	EVVREVCEGAETGECKALLESKWIFFUNGER DEFEYGEGGGRENNEDTEEVCHAVGGAI	61.00		3641.0	288.0
TW6167	EVVREVCERIALENCE CONTRACTOR OF THE CONTRACTOR OF	34.00			
BG031	EVVREVCSELAEVAR CHARLES OF THE TOTAL STATE OF THE STATE O	49.00		731.0	
BG032	EVYREVCSE ABSOLLOME SOM SENTENCE OF STATEMENT OF STATEMEN	2000.00	11.50		
TW101	BVCBEAMINE CHARLES OF THE CONTRACTOR OF THE CHARLES			369.0	
TW6208	BUNREYCE BLAB TAF CHARLES OF THE STATE OF TH	260.00	3.70		
TW106	EVESEGABLIOTE CANALISATION OF THE STATE OF T	1.70	11.20	1600.0	123.0
00108	EVVREVCSELATION TO SERVETANT BOLD PET Y GGCGGURAN FOT BEYCHAVCGSAI	9.50		1681.0	421.0
00100	EVVRBV, BELANTION CONTINUED TO SERVICE STATEMY PROTECTION OF SERVICES AT THE SERVICES AT THE SERVICE STATEMY PROTECTION OF SERVICES AT THE SERVI	2.10		624.0	65.0
DD110	KVVKEVCSEGAB LOTCH STATEMENT BOX OF PROCESSING STATEMENT BOX ON A STATEMENT BOX OF STATEMEN	5.60			
DD111	BVVKEVUS MARIOTORS 121 TO BY THE TO BE PYCHOLOGICAL TO BE TO BE THE TOTAL TO B	9.90		998.0	
DD112	EVVREVCSELARITOFCHAY ISAMINEDITE OF THE STATE	78.00		368.0	
TW6179	EVVRKVCS BLANTIST MALES AND SELVEN PROPERTY PER Y GOOGGINEN POTERY CHAVCES A I	4.70	103.58	4532.0	457.0
TW6163	EVVREVCSELAR I GENERALISM GENERAL	315.00			1463.0
TW6172	EVAREVCESE/ARTIGECEALING TO THE TOTAL TOTA	70.00		885.0	39.0
TW6180	EVVREVUBIGATE TO THE THE CONTRACT THE THE THE THE THE THE THE THE THE TH	150.00		1514.0	
TW6181	EVVERUCEDIMETER CONTROLL SPANFORM CAPPYORCE APPYORCE CHANGES AND CONTROLL SPANFORM CAPPYORCE AND C	38.00	10.00	489.0	204.0
BG001	EVYREVC SELMETOT TOWN TOWN TOWN TOWN TOWN TOWN TOWN T	145.00	00.69		0.908
TW118	BVCSGJABITOFCHARALIAN I CONTROLLED BY CGCCGGURANPDTEEYCHAVGGSAI	16.00		315.0	
20102	BUVREVUSELING TO THE TRANSPORT OF THE TR	17.00	0	2128.0	110.0
DD103	KVVKEV SELPELOF COMPANY TO THE SELECT SEPTION OF THE SELECT SELEC	15.00	0	237.0	345.0
DD104	KVVKKVLSKLMBIOTCHOWNING THE TOTAL APPRYCECOGNINNFOTEETCHAVGGSAI	18.00		198.0	320.0
50106	EVVKEVCSELENDISECTORISE	25.80	0	3521.0	395.0

EIG 16/0)	_	Inhibition Ki (nM)	Ki (uM)	
77/04 - 012	P. kall	Plasmin		%
DVTEGKCAPFYGGCGGNRNNFDTEEYCMAVCGSAI	36.00		762.0	
WTEGECAPFFYGGCGGNRANFDTEEYCMAVGSAI	70.83			
WTEGKCAPFFYGGCGGNRAINFDTEEYCMAVCGSAI	54.00		277.0	
WTEGKCAPFFYGGCGGNRNNFDTEEYCMAVCGSAI	110.20		0.00968	133.0
WTECKCAPPFYGCCGGNRNNFDTEEYCMAVCGSAI			40.0	116.0
NTEGRCAPFYGGCGGNRNNFDTBBYCHAVCGSAI	81.00	45.90	184.0	613.0
DVTEGKCAPFFYGGCGGNRNNFDFBBYCHAVCGSAI	184.00		402.0	
DYTECKCAPPYGCCGCNRNNFDTEBYCMAVCGSAI	44.00			37.0
TATEGREAP FUY GOOGGINENIN POTERY CHAVEGS A I	18.00	18.00	7972.0	225.0
DYTEGECAPFLYGGCGGNRNNPDTEEYCNAVCGSAI	216.00		1557.0	
DVTEGKCAPFGYGGCGGNRNNPDTEEYCHAVCGSAI	39 00			316.0
DVTEGKCAPPTYGGCAGNRNNPDTEEYCHAVCGSAI	35.00		1090.0	179.0
DYTECKCAPPYGGCKGNRNNFDTEEYCHAVCGSAI	18.00		921.0	309.0
DYTECKCAPFYGGCLGNRNNFDTEEYCHAVCGSAI	11.00		915.0	39.0
DYTECKCAPPYGCCHGNRANFDTEEYCHAVCGSAI	11.00			27.0
DATEGICAPPYGGCNGNRNNFDTEEYCHAVCGSAI	35.00		475.0	
DVTECKCAPPYGGCPGNRNNFDTEEYCMAVCGSAI				
DVTEGKCAPFYGGCQGNRNNFDTEBYCHAVCGSAI	42.00			
DVTBGKCAPFFYGGCRGNRNNFDTEEYCHAVCGSAI	6.00	24.00	13009.0	0.89
DVTBGKCAPFYGGCCGNRNNFDTEEYCMAVCGSAI	15.00			
TOVTECKCAPFYGGCSGNRNNFDTBEYCHAVCGSAI	40.00		511.0	168.0
DVTEGKCAPFFYGGCTGNRANFDTEEYCMAVCGSAI	29.00			
DVTEGKCAPPYGGCVGNRNNFDTEEYCMAVCGSAI	17.00			0.4.0
PWTEGRCA PPY GGCYGNRANPDTEEYCHAVCGSA I	7.50	18.00	1507.0	8.7
POVTEGRCAPPYGGCDGNRNNFDTEEYCHAVCGSAI	64.00		924.0	
PUTEGRCAPFYGGCEGNRNNPDTEEYCMAVCGSAI	163.00		1162.0	954.0

Inhibition Ki (nM)

		16,40(3)	i	i	3	\$
	Variant	Sequence	8	Plasmo		8
		SOUTH STANSFORM SOUTH SOUTH STANSFORM SOUTH SOUTH STANSFORM SOUTH STANSFORM SOUTH STANSFORM SOUTH STANSFORM SOUTH STANSFORM SOUTH SOUTH STANSFORM SOUTH STANSFORM SOUTH STANSFORM SOUTH STANSFORM SOUTH STANSFORM SOUTH SOUTH STANSFORM SOUTH STANSFORM SOUTH STANSFORM SOUTH STANSFORM SOUTH SOUTH STANSFORM SOUTH STANSFORM SOUTH STANSFORM SOUTH SOUT	19.00	22.80	152.0	78.0
	TW6139	KVVRKVCSKLAKISTOTOMATISMATISMATISMATISMATISMANIFITRRVCMAVCGSAI	11.20	21.30	65.0	36.0
	TW6153	EVVREVCSEQAETGFCKAALSKWIF DVIEGRAFF I 1000 LOMBER 1000	32.00	27.00		581.0
	TW122	EVCSEQAETGPCRANISRNYPDVTEANCAPT FILACCAMMUM DISELECTION OF THE STATE O	16.00		444.0	
	TW6178	EVVREVCSEQAETGPCRAMISHWI PUVIDAN. AFF FIGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	40.00			
	TW8148	BVVREVCSBOAFTGPCRAMISKWYFLVTEGACAFFF I USCUSARAMIE DI ESTERNICO EN TENTONICO EN TEN	64.00	48.00		
	TW124	KVCSEQAETGPCRAMISRMYFIVTEGKCAPFFYGGCGGNSNNFUTEETCHAVCGSA1	8			
S	1	EVVREVCSEQAETGPCRAMI SRMYPDVTEGRCAPPPYGCCGGNANNPDTEEYCMAVCGSAI	3.5			T
UI	1	PRIN BUCSPOA BTGPCRAAHSTNYPDVTBGKCAPPYGGCGGNRNNPDTBEYCMAVCGSAI	1.00	7.24	1432.0	
35	ı	THE PROPERTY HARMY FINTEGRICA PETYGGGGGNRINIF DTEETCHAVCGSAI	0.90	6.89	2796.0	
П	TW6174	EVANEVOSE, MEDICINE STATEMENT POR CAPPY OCCOUNTINFOREY CHAVCOSAI	0.98	19.00	403.0	0.09
TU	BG002	EVVREVCSKIAKTAKTAKTAKTAKTAKTAKTAKTAKTAKTAKTAKTAKTA	3.60		1864.0	6.0
TE	DD129	EVVREVCSEQAETORCKALFSKWIFDV IEGALAFF I I I I I I I I I I I I I I I I I I	0.39	17.8	150.0	196.0
5	DD185	EVVREVCSEQAETGPCRAALFRMIFDVIEGRCAFFILGCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	0.65	16.40	206.0	
HE	TW6165	BUVREVCSBQARTGPCRAAIWRWYFDVTBGACAFFF IGGCGGARANIC CONTROLL	940	10.10	73.0	
Ε		EWREVCSEQAETGPCRAAIYRMYFDVTEGRCAPFFYGGCGGNRNNFDFEETCRAVCGSAL			9 60	
T (•	RAVARIVES BOART OPERALIY RWY FOUTBOICA PFY GGCGGNRNN POTERY CHAVCGSAI	2.7	12.10	62.0	
RI		THE STATE OF THE PROPERTY OF T	1.20		619.0	111.0
JL	•	EVVREVCEDAR I DE CACAL TE DE MY PINTEGECA PPFYGGCGGNRNNFDTEEYCMAVCGSA I	0.85	12.80	293.0	74.0
E 2	'	EVVREVC SELATE TOTAL TERMY PRAMERICA PEPY COCCOURANTETTE CHAVES A I	0.50	7.46	35.0	99.0
6)		EVVREVCSBUARTISECTION OF THE STATE OF THE ST	34.60		419.0	
	TW8201	EVURBUL SELICIONES CONTRACTOR DE PROCESCANDAMENTE EXCHAVOSA I	128.50		1237.0	
	TW6202	EVVREVCEDQAKTGFCKAGIMKMI FDV I EGINCALI I SOCIONALINE STOREN SELECTION DE STOREN SELECTION SELEC	31.20		5045.0	
	TW6203	EVUREVCSEQAETGPCRAGI PRINTE DVI EGACARTE E I DOCCOMINIONE DE LOCALINA DE LA COMPANIONE DEL COMPANIONE DE LA COMPANIONE DEL COMPANIONE DE			147.0	87.0
	TW6204	EVVREVCSEQAETGFCRAALSAMI FUVIESANAFF FISCOCATION BETWEEN CHANGES I			196.0	29.0
	TW6205	EVVREVCSEQAETGPCRALI SAWYFDVTESICAPFFICACCAMMAN DI BEI CHANCESIN	0,0	1111	224.0	
	DD114	EVVREVCSEQAETGPCRAAISRMYFDVTEGKCAPPFYGGCRGNRNNFDTEETCHAVCUSAL	2 8		0 084	1398.0
	TW6190	RVVREVCSEQAETGPCRAAISRMYFDVTEGKCAPFFYGGCYGNENNFDTEEYCMAVCGSAI	20.05	_	1	469.
	TAKETES	RVVREVCSEOAETGPCRAMHPRWYFDVTEGKCAPPFYGGCGGNRNNPDTEEYCMAVCGSAI	1.20	11.00	12440.0	2.62
	Moios					

Inhibition Ki (nM)

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46
<u>6</u>
سا

Sequence	ਹ	E kall	Plasmo	4	A
CALLE DESCRIPTION OF AMERICAN PROCESSOR CALLE DESCRIPTION OF THE CONTROL OF T	SAI	0.91	11.96	14000.0	214.0
SATISTICS OF STATES OF SATISTICS OF SATISTIC	SAI	1.30	18.60	388.0	473.0
BAVARACISE AND HEAVED THE PROPERTY OF THE STATE OF THE ST	-	38.00		467.0	
ENVERTISED FOR THE PARTICIAL PROPERTY OF THE P	SAI	0.48	9.96	186.0	11.0
EN ARENCERA RECOGNAMI PRINT PONTECKCA PPFYOGCYGNRUNP DFEEVCHA VCGSA I	SAI	3.80	15.40	92.0	16.0
SAMPHANTSBOAKTOPCRAMIYPRY POVTBOKCA PFFY GGCY GURINNF DTEEY CMAVCGSA I	SSAI	8		419.0	24.0
ENTRE HANDEN CRAMINEN Y POYTECK CAPETY OCCYCURINED TESY CHAVCOSAI	3SAI	8.4			34.0
EVVIEW CENTRAL SECRET SERVED SECONDARY SECONDARY SECONDARY SECONDARY SERVED SER	GSAI	2.50			452.0
ENAME EN A PROPERCY I TRINY FIVT EGICA PPY GCCCCONTRINY FOR ECCENT	GSAI			213.0	289.0
PHATA PHATA PROPERATION FOR TECK CAPPY COCOCARANT DE LEY CHAVEGSAI	GSAI	0.09	18.00	550.0	
DATE OF THE PROPERTY PRINTEGE CAPETY GOOD OF THE PROPERTY OF T	GSAI	3.50	118.00	56.0	
EVANEWOCSEGA ETGPCRALHPRIVY FDVTEGKCA PFFYGGCGGNRINI FDTEEYCHAVCGSA I	GSAI	7.20	32.70	245.0	156.0
ENVIRONSEDAETGPCRAALFRHYPDVTEGRCAPFFYGGCGGNRNNFDTEEYCNAVCGSAI	GSAI	0.30	12.10	80.0	
PAALBENCEDA ETGPCBA L. PTRIVY POVTEGECA PFYGGCGGNRUNF DTEEY CHAVCGSA I	GSAI	6.60			9.5
WANDENCEDA ETGPCRALFKRMY FONTEGECAP F PYGGCGGNRNN F DTBBY CHAVCGSAI	GSAI	7.90	2.00	1385.0	3.3
WIND BY CROAPTOPCRAPPIRMY PONTEGRICA PPY OCCOURANT POTERY CHAVCGSAI		112.00			16.8
NAME OF THE PROPERTY OF THE OWNER OF THE OWNER OF THE OWNER OF THE OWNER	CSAI	9.30			11.0
RAVE ENCROAET OPCRALLS ANY FOUTEGRICA PFY GGCGONRIN POTERY CHAVCGSA I	GSAI	19.00			21.0
EVVR EVCSEOAETGPCRALIMHWYFDVTEGKCAPPYGGCGGURINIPDTEEYCHAVCGSAI	GSAI	9.20	18.70	18.0	
EVVR EVCSBOARTOPCRALI PANYPDVTBGKCA PPPYGGCGGARNNPDTEEYCHAVCGSA I	GSAI	15.00			46.0
EMARENCEDA ETGPCRALI YHMYPDVTEGKCA PPYGGCGGARNNFDTEEYCHA VCGSA I	GSAI	6.00	12.20	19.4	697.0
MADENCEDA RECOCRAA I HKWY PDVTEGKCA PFYGGCGGNRNN FDT REY CHAVCGSA I	GSAI	1.70		106.0	
EVVREVCSEOAETGPCRAAIYHMYPDVTEGRCAPPYGGCGGRBANPDTEEHCHAVCGSAI	GEAI	0.64	7.26	14.5	
EVVREVICEBOAETQPCRALIQHMYPDVTBGRCAPFFYGGCGGRRNNPDTEEYCHAVCGSAI	CGSAI	23.00		262.0	
ENVREVISEOAETGPCRALIYIONYPDVTBGKCAPPYQGCGGURNNPDTEEYCMAVCGSAI	CGSAI	4.10	7.47	38.7	
WALD EVEN SEDA ETGPCRAA I OHEN PIDVTEGKCA P PY OCCOGNENNIP DTEEY CHAVCGSAI	CGSAI	5.80		144.0	

TW618TW

	FIG. 46(5)	2 2 3	Inhibition Ki (nM)	Ki (nM)	晃
Variant	Sequence	0.14		583.0	84.0
DD118	EVVREVCSBQAETGECKAALFERFIED LONCAFFF LOCKONFINEDCESCHAVOSAL	0.26		0.499	20.0
TW6191	BVVREVCEBOARTOPCKAALFKAFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	0.11		1034.0	99.0
00117	EVVREVCSEQAELIGECKALLERANIE CONTRACTOR EN PROCESSION EN PROCESSA E	3.20		7.9	
80029	EVVKEVCSEGREVSECTORS. EVVKEVCSEGREVSECTORS. TVHYPDVTEGKCA PFYGGCGGRRINFDTEFYCHAVCGSA I	4.60		26.1	
BG030	BVALEVOSEMINOS CONTROLLOS SENTINOS DE PROCESORIAN ED TERICANA VEGASA I	0.75		5.8	
BG033	EVALENCE CANDESCRIAL I YHMY FDVTEGKCAPFYGGCGGNRUNPDTEEYCHAVCGGAI	0.47		18.5	
BGG34	RAVREVCSBOARIGPCRALIYHMYPDVTBGRCAPPPYGGCGGRRANPDTEEYCMAVCGSAI	3.40		9.6	
90046	EVVREVCSEQAETGPCRGAIQHMYFDVTEGKCAPFPYGGCGGNRANPDTEEYCHAVCGSAI	160.00		178.0	
2 :	EVVREVCSEOAETGPCRGAIRHWYFDVTBGKCAPFFYGGCGGNRNNFDTEEYCMAVCGSAI	- 80.08		200.0	
BGUI	PRINGENCEDAR PROPURES I RHMYPDVT BGKCAP PYGGCGGNRUNPDYEEYCMAVCGSAI	340.00		224.0	
BG021	BY VIEW CONTROL OF THE POST BOX CAPPING COGNINAL DIERY CHAVCESAI	65.00		18.2	
BGGZP	EVALUACIONO PRODUDOS I VANA PINTEGICA PPYGGCGGARIAN FOT BEYCHAVICES A I	50.00		34.9	
BG026	EVVREVCERGALISTOCKA TIMONVERGECA PPPYGGCRIQARANIEDTEEVCHAVCGSAI	0.53			
DD118	EVVREVE BEGAB I OF CHARLESTON OF THE STATE O	1.10	1.05	15640.0	9.0
00134	BVVKKVCSEJALIGICKARIE KANILIS OF TOTAL CARRANDOTTERYCKAVCGSAI	1.30		7473.0	0.0
DD136	EVOREVCSEQAETGPCKALFAMITED TECHNICATION CONTRIBUTED OF CHANGES AT	100			1.8
200	EVVREVCSEQAETGPCRALFERMYPDVTEGACAPFFTGG, FRANKUNFD1BB1CHNVCGGA				

FIG. 47

VOLUM	AES	
NS	344.25	
KPI	245.75	
	KPI N	S
	298	366
	266	342
	354	294
	258	385
	168	288
	266	469
	172	338
	184	272
MEAN	245.75	344.25
STDEV	66.2414415	63.97488346
TTEST		0.009094999
	-	

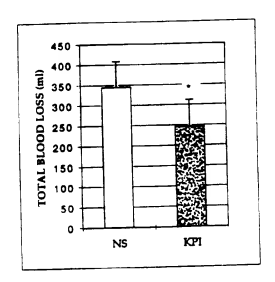


FIG. 48

TIENCO	TI OPINI	
HEMO	GLOBIN	
NS	23.61	
KPI	13.59	
	KP1	NS
	16.58	24.95
	15.19	24.87
	20.21	20.46
	8.99	27.59
	14.63	18.23
	15.31	31.59
	7.7	23.26
	10.14	17.96
MEAN	13.59375	23.61375
STDEV	4.261438	4.68761
TTEST		0.000536

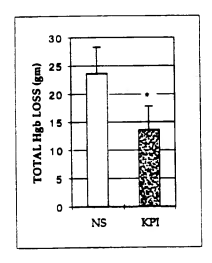


FIG. 49

	Baseline PaO2	Pa02	End CPB	CPB		Obs 60 min	nin	<u>5</u>	bs 180 min	nir
	KP	9	Ē	2	NS NS	Ē	SN	Z.		NS
	652.2	6029	•	195.7	60.5	483.7	441.3	_		391.3
	35	569.2	_	974	132.2	330.1	448.7		264.1	484.6
	5.96.2	6229	_	70.2	8.68	415.4	1 85.1		416.5	81.3
	606.2	689.2	_	264.2	333.9	430.2	529.6		361.9	333.2
	633.1	665.1	-	67.2	341.7	613	568.3	_	8.08	546.6
	446	527		07.4	226.9	564.3	438.1	_	518.2	485.3
	563.2	461.7		47.1	89.1	<u> </u>	42.6	<u>.</u>	494.2	45.6
	6 659	Ş	_	416.6	59.7	504.5	8.904	-	452	383.7
Z	626.425	88	426	426.625	167.225	480.275	369.938	L_	371.1	344
D.F.V	24 46973	85.50556	140	140.4741	117.9931	88.6187	88.61879 196.5235		150.2774	186.227
EST	12	0.268		=	0.0014	=d	0.17915		<u>_</u>	0.76
		914					N.S.			

.02

	F16. 50		Summary of Data	ny of	Data		•		-
			Total Volumes	mes	0.7	Serial Ch	Serial Chest tube Hbg	Hbg	
	Total volume loss	Total Hgb Loss	Chest tube Sacrifice	Sacrifice		0-30min	30-60min	60-120min	30-60min 60-120min 120-180min
10.2	$\overline{}$	16.58	8	113		3.7	4.3	8.6	6.2
1-1-2	776	15.19	198	88	1	4.3	6.4	6.7	5.7
KPI-2	730	20.21	12	212	·	4.1	4.4	7	7.1
KPI-3	250	8	5	88		2.8	4	4.4	1.9
KPI 4	907	14 63	8	2		6.3	6.5	7	6.7
KPI-5	168	15.31	188	78		4.1	6.1	5.6	6.3
KPI-6	8 2	7.7	134	38		3.1	4.6	5.4	4.4
X Z	7/1	41.01	158	26		6.9	5.8	5.4	4.2
KPI-8	132	10.11			_				
	200	13.50			MEAN	4.41	5.26	6.26	5.3
MEAN	245.75	2001			STDEV	1.45	1.04	1.32	1.72
STDEV	66.24	4.20							
				8		7.7	8 6	19	5.4
NS-1A	366	24.95	274	7					
NS-2	342	24.87	236	<u>5</u>		7.7			
NG.	767	20.46	757	42		5.4			
NG A	385	27.59	303	82		8.4	7.2		
2 2	288	18.23	140	148		7.5	7.2		
2 2	469	31.59	197	208					
NG 7	338	23.26	218	120		7.5		7 5.8	
85Z	22	17.96	902	%		7.4	8.2		6 5.3
					1	08.7	76	A 58	61
MEAN	344.25	23.61			MRAN	0.0	1		
STOEV		4.69			STDEV	1.44	1.04	16.0	U.83
	% = 0.009	$^{\bullet}p = 0.0005$				\$00.0 = d	"p = 0.004 "p = 0002	NS	NS

pTW 6166

FIG. 51

α-factor ATG AGA TIT CCT TCA ATT TIT ACT GCA GTT TTA TTC GCA GCA TCC TCC GCA TTA GCT TAC TCT AAA GGA AGT TAA AAA TGA CGT CAA AAT AAG CGT CGT AGG AGG CGT AAT CGA Met Arg Phe Pro Ser ile Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG, TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG ≯Ala Pro Val Asn Thr Thr Glu Asp Glu Thr Ala Gln II. Pro Ala Glu Ala Val ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG File Gly Tyr Leu Asp Leu Glu Gly Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn AGC ACA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TCT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT Ser Thr Asn Asn Gly Leu Leu Phe Ile Asn Thr Thr Ile Ala Ser Ile Ala Ala Lys KPI(-4-57; M15A, S17Y) GAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA Xbal CTT CTT CCC CAT AGA GAT CTA TTT TCT CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT ▶Giu Giu Giy Val Ser Lau Asp Lys Arg Giu Val Val Arg Giu Val Cys Ser Giu Gin RsrII Aatll Agel SCT GAG ACC GGT CCG TGC CGT GCA GCT ATC TAC CGC TGG TAC TTT GAC GTC ACT GAA CGG CTC TGG CCA GGC ACG CCA CGT CGA TAG ATG GCG ACC ATG AAA CTG CAG TGA CTT Ala Glu Thr Gly Pro Cys Arg Ala Ala IIe Tyr Arg Trp Tyr Phe Asp Val Thr Glu GGT SAG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC AAC CGT AAC AAC TTT GAC CCA TTC ACG CGA GGT AAG AAA ATG CCG CCA ACG CCG CCG TTG GCA TTG TTG AAA CTG FGIY Lys Cys Ala Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp BamHI ACT GAA GAG TAC TOC ATG GCA GTG TGC GGA TCC GCT ATT TAA GCT T TGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A FThr Glu Glu Tyr Cys Met Ala Val Cys Gly Ser Ala 11e

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α-factor
ATG AGA TIT CCT TCA ATT TIT ACT GCA GTT TTA TTC GCA GCA TCC TCC GCA TTA GCT
 TAC TOT AMA GGA AGT TAM AMA TGA CGT CAM MAT AMG CGT CGT AGG AGG CGT MAT CGA
Met Arg Phe Pro Ser IIe Phe Thr Ala Val Leu Phe Ala Ala Ser Ser Ala Leu Ala
 GCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG
Ala Pro Val Asn Thr Thr Glu Asp Glu Thr Ala Gln Ile Pro Ala Glu Ala Val
 ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC
 TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG
▶!!e Gly Tyr Leu Asp Leu Glu Gly Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn
 AGC ACA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT
Ser Thr Asn Asn Gly Leu Leu Phe ile Asn Thr Thr lie Ala Ser ile Ala Ala Lys
                                           KPI(-4-57; M15L, S17F)
                      Xbal
 ▶Giu Giu Giy Vai Ser Leu Asp Lys Arg Giu Vai Vai Arg Giu Vai Cys Ser Giu Gin
             Rsrll
                                                                      Aatli
          Agei
 GCT GAG ACC GGT CCG TGC CGT GCA TTG ATC TTC CGC TGG TAC TTT GAC GTC ACT GAA CGA CTC TGG CCA GGC ACG GCA CGT AAC TAG AAG GCG ACC ATG AAA CTG CAG TGA CTT
PAIR Glu Thr Gly Pro Cys Arg Ala Leu Ile Phe Arg Trp Tyr Phe Asp Val Thr Glu
GGT AAG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC AAC CGT AAC AAC TTT GAC CCA TTC ACG CGA GGT AAG AAA ATG CCG CCA ACG CCG CCG TTG GCA TTG TTG AAA CTG
FGIY Lys Cys Ala Pro Phe Phe Tyr Giy Giy Cys Giy Giy Asn Arg Asn Asn Phe Asp
                                           BamHI
 ACT GAA GAG THE TOO ATG GEA GTG TOO GGA TOO GCT ATT TAA GCT T
 TGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A
>Thr Giu Giu Tyr Cys Met Als Val Cys Gly Ser Als lie
```

FIG. 53

α-factor ATG AGA TIT CCT TOA ATT TIT ACT GOA GIT TIA TIC GOA GOA TOO TOO GOA TIA GOT TAC TOT AAA GGA AGT TAA AAA TGA CGT CAA AAT AAG CGT CGT AGG AGG CGT AAT CGA Met Arg Phe Pro Ser ile Phe Thr Ala Vai Leu Phe Ala Ala Ser Ser Ala Leu Ala CCT CCA GTC AAC ACT ACA ACA GAA GAT GAA ACG GCA CAA ATT CCG GCT GAA GCT GTC CGA GGT CAG TTG TGA TGT TGT CTT CTA CTT TGC CGT GTT TAA GGC CGA CTT CGA CAG PAIA Pro Val Asn Thr Thr Glu Asp Glu Thr Ala Gin ile Pro Ala Giu Ala Vai ATC GGT TAC TTA GAT TTA GAA GGG GAT TTC GAT GTT GCT GTT TTG CCA TTT TCC AAC TAG CCA ATG AAT CTA AAT CTT CCC CTA AAG CTA CAA CGA CAA AAC GGT AAA AGG TTG File Gly Tyr Leu Asp Leu Glu Gly Asp Phe Asp Val Ala Val Leu Pro Phe Ser Asn AGC ACA AAT AAC GGG TTA TTG TTT ATA AAT ACT ACT ATT GCC AGC ATT GCT GCT AAA TCG TGT TTA TTG CCC AAT AAC AAA TAT TTA TGA TGA TAA CGG TCG TAA CGA CGA TTT Ser Thr Asn Asn Gly Leu Leu Phe Ite Asn Thr Thr Ite Ala Ser Ite Ala Ala Lys KPI(-4-57; M15L, S17Y) Xbal GAA GAA GGG GTA TCT CTA GAT AAA AGA GAG GTT GTT AGA GAG GTG TGC TCT GAA CAA CTT CTT CCC CAT AGA GAT CTA TTT TCT CTC CAA CAA TCT CTC CAC ACG AGA CTT GTT ⊳Giu Giu Giy Vai Ser Leu Asp Lys Arg Giu Vai Vai Arg Giu Vai Cys Ser Giu Gin RsrII Aatii Agel GCT GAG ACC GGT CCG TGC CGT GCA TTG ATC TAC CGC TGG TAC TTT GAC GTC ACT GAA CGA CTC TGG CCA GGC ACG GCA CGT AAC TAG ATG GCG ACC ATG AAA CTG CAG TGA CTT PAIR GIU Thr Gly Pro Cys Arg Ala Leu lie Tyr Arg Trp Tyr Phe Asp Val Thr Glu GOT ANG TGC GCT CCA TTC TTT TAC GGC GGT TGC GGC GGC AAC CGT AAC AAC TTT GAC CCA TTC ACG CGA GGT AAG AAA ATG CCG CCA ACG CCG CTG GCA TTG TTG AAA CTG FGIV Lys Cys Ala Pro Phe Phe Tyr Gly Gly Cys Gly Gly Asn Arg Asn Asn Phe Asp BamHI HindIII ACT GAA GAG TAC TOC ATG OCA GTG TGC GGA TCC GCT ATT TAA GCT T TGA CTT CTC ATG ACG TAC CGT CAC ACG CCT AGG CGA TAA ATT CGA A

>Thr Glu Glu Tyr Cys Met Ala Val Cys Gly Ser Ala !!e

FIG. 54(1)

PROTEIN	BONDACE	K. kallikrein	K, Factor XIIa	E. Plesain
	COUNTRY CONTRACTOR	22.6	2000	0.33
Aprotinin	RPOFCLEPPTTOPCKARIIRTFTNAKAGLCOITVIGGCKANGNE NOALDGELEE	45.0	3718.0	34.00
KP1(-4-57)	EVVIEWCSEQAETGPCHAMISRAT PDVTEGECAPPT TGCCGGGNNAR DIEBLICHANGGSAI	>5000	pu	12.30
TW101	EVVREVCSEQNETGPCKANISRWY FDVTEGKCAPTFTGCCGGGRRWW DIESTCHWYCGSAI	449.0	pu	2.98
TW106	EVVREVCSEQAETGPCHARISHWY PDVTEGKCAPTFTGGCGGGRNWRF DIE ENGENIE	116.00	þu	70.90
TW116	EVVREVCSEQAETGPCRANIIBATFDVTEGKCAPFTGGCGGRRRR DIE ELCHINGE	>5000	þu	1.45
TW105	EVVREVCSEQAETGPCEARISHAY FDVTEGKCAPT FTGGCCGGNUNG F D SECTION	>\$000	PE	19.90
THII7	EVVREVCSEQAETGPCKANI I RWYFDVT EGKCAPFFYGGCGGNNNN FOTEET CHAVCUSAL	631.0	þ	2.24
TH115	EVVREVCSEQAETGPCRAELIRMYFDVTEGRCAPFFYGGCGGNRNNFOTEETCHAVCUSAL	25000	PE	1.27
TW102	EVVREVCSEQAETGPCKARIIRWYFDVTEGKCAPTFYGGCGGNANNFOTEETCHAVCSAL	000	>5000	>5000
CL005	EVVREVCSEGAETGPCAMISBWY FDVTEGKCAPFFYGGCGGNRNNFDTEEYCHAVCGSAI	236.0	9	1555.0
TH6172	EVVREVCEEGAETGPCRAKASRWY FOVTEGKCAPPTYGGGGGNRNNFDTEETGAAVGSAL	21515	615.0	44.10
TW6207	EVVREVCSEQAETGPCAMIABMY FDVTBGKCAPFFTGGCGGMANN FDTERCHAVCOSAL	110.2	00969	31.10
CT-0062	EVVREVCSEQAETGPCRANTSANYFDVTBGKCAPFFTGGCGGHKNNFDIEELCKAVCGGA	1.7	1600.0	11.20
DD108	EVVREVCSEQAETOPCHALISMY FDVTECKCAPF FT CCCCCGNNN FD LEEL CON CONTROL		624.0	11.000
00110	EVAREVESEQAETGPCRALISANTEDVIEGKCAPFFTGCCGGNKNNFDIEELCKAVCGSTI	,	Pu	Pu
DD111	BUNNEVCSEQNETGPCRASISEMY FDUTEGKCAPFFFGCCGGNRNNFDTEETCAN CUSAL	9	0.866	P
DD112	EVVREVCSEQAETGPCRAVISENT FDVT EGKCAPF FT GGCGGNRNN FDT EET CHAVCGSAT	2.5	0.511	9
00102	EVVREVCSEQAETGPCRAHIPRWY FDVTEGKCAPFFYGGCGGNRNNFDTEET CHAVCGSAI	0.91		5
00103	EVVREVCSEQAETGPCRAHIFRWYFDVTEGXCAPFFYGGCGGNRNNFDTEETCHAVCGSAI	0.11		1 2
DD104	EVARVESEGAETGPCRAHITRWYFDVTEGKCAPFFTGGCGGNRNNFDTEETCHAVCGSAI	13.0	2	2 2
00100	EVVREVCSEQAETGPCBAHIARMY FDVTEGKCAPFFYGGCGGNRNN FDTEEY CHAVCGSAI	0.81	33.0	10. 10
TW6166	EVVIEVOSEGAETGPCKAALLEMY PDVTEGKCAP F T T OG COGNEMN F DT T ET CHAVCOSA I	• •	2121	

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FIG. 54(2)

		6.5	206.0 16.4	16.4
	EVVREVCSEQAETGECKAA MANIE DVI LOKAKEV TOGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG			
	PROPERTY SPONTED TO BE TO BE TO BE THE SECOND STATES OF THE SECOND STATE	1.1	93.8 12.10	12.10
			15.0 7.46	7.46
1=6	EVVREVCSEQAETGPCNAL ITAMY FDVTEGKCAFFFTGGCGGHNNFDIELICNAVCGSAI	6:5		
, , ,	PHYSEVOSECAPTGPCRANIMMYPDYTEGECAPPYGGGGGHANNPOTEETCHAVGGSAI 2.5	2.5	40.0 nd	pu
		6.6	9/	7
1×62 %	EVVREVCSEQUEIGE CROAD HOMIT DA I L'ANCART FINANCHER ELECTRICE			
TW624	EVYREVCEBOALTOPCHAALHENT PDVTEGECAPPTT000080000FDTESTCHAVC8SAI 4.6	4.6	36	T

